PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7: C12N 15/82, C07K 14/80, A01H 5/00

A2

(11) International Publication Number:

WO 00/09720

(43) International Publication Date:

24 February 2000 (24.02.00)

(21) International Application Number:

PCT/GB99/02676

(22) International Filing Date:

13 August 1999 (13.08.99)

(30) Priority Data:

PP 5268

14 August 1998 (14.08.98) AU

(71) Applicants (for all designated States except US): INTERNATIONAL FLOWER DEVELOPMENTS PTY LTD [AU/AU]; 16 Gipps Street, Collingwood, VIC 3066 (AU). VRIJE UNIVERSITEIT [NL/NL]; De Boelelaan 1087, NL-108 HV Amsterdam (NL).

(71) Applicant (for GB only): JONES, Elizabeth [GB/GB]; Frank B. Dehn & Co., 179 Queen Victoria Street, London EC4V 4EL (GB).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): KOES, Ronald [NL/NL]; De Boelelaan 1087, NL-108 HV Amsterdam (NL). DE VETTEN, Nick [NL/NL]; De Boelelaan 1087, NL-108 HV Amsterdam (NL). MOL, Jos [NL/NL]; De Boelelaan 1087, NL-108 HV Amsterdam (NL).
- (74) Agents: JONES, Elizabeth et al.; Frank B. Dehn & Co., 179 Queen Victoria Street, London EC4V 4EL (GB).

(81) Designated States: AE, AL, AM, AT, AT (Utility model), AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, CZ (Utility model), DE, DE (Utility model), DK, DK (Utility model), DM, EE, EE (Utility model), ES, FI, FI (Utility model), GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK (Utility model), SL, TJ, TM, TT, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European

Published

Without international search report and to be republished upon receipt of that report.

patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF,

CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

(54) Title: CYTOCHROME B5 FROM PETUNIA

(57) Abstract

The present invention relates generally to a plant regulatory gene and derivatives and homologues thereof. More particularly, the present invention provides a nucleic acid molecule comprising a sequence of nucleotides or complementary sequence of nucleotides, the expression of which, modulates or otherwise facilitates activity of a cytochrome P450 protein in plant cells and tissues including petals, flowers, stems, leaves and seeds. Even more particularly, the nucleic acid molecule of the present invention encodes a cytochrome b₅ or a mutant, part, fragment, portion, functional and/or structural equivalent or homologue thereof or agonist or antagonist thereof involved in modulating or otherwise facilitating activity of a dihydrokaempferol (DHK) hydroxylating enzyme such as but not limited to flavonoid 3', 5'-hydroxylase. The present invention further provides transgenic plants or parts thereof or cells of transgenic plants as well as cut or severed flowers or stems from transgenic plants.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland ,	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	\mathbf{SZ}	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	$\mathbf{U}\mathbf{Z}$	Uzbekistan
CF	Central African Republic	JP	Japan -	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Suđan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

A PLANT REGULATORY GENE

The present invention relates generally to a plant regulatory gene and derivatives and homologues thereof. More particularly, the present invention provides a nucleic acid molecule comprising a sequence of nucleotides or complementary sequence of nucleotides, the expression of which, modulates or otherwise facilitates activity of a cytochrome P450 protein in plant cells and tissues including petals, flowers, stems, leaves and seeds. Even more particularly, the nucleic acid molecule of the present invention encodes a cytochrome b₅ or a mutant, part, fragment, portion, functional and/or structural equivalent or homologue thereof or agonist or antagonist thereof involved in modulating or otherwise facilitating activity of a dihydrokaempferol (DHK) hydroxylating enzyme such as but not limited to flavonoid 3', 5'-hydroxylase. The present invention further provides transgenic plants or parts thereof or cells of transgenic plants as well as cut or severed flowers or stems from transgenic plants.

15

Throughout this specification, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element or integer or group of elements or integers but not the exclusion of any other element or integer or group of elements or integers.

20

The subject specification contains nucleotide and amino acid sequence information prepared using the programme PatentIn Version 2.0, presented herein after the bibliography. Each nucleotide or amino acid sequence is identified in the sequence listing by the numeric indicator <210> followed by the sequence identifier (e.g. <210>1, <210>2, etc). The length, type of sequence (DNA, protein (PRT), etc) and source organism for each nucleotide or amino acid sequence are indicated by information provided in the numeric indicator fields <211>, <212> and <213>, respectively. Nucleotide and amino acid sequences referred to in the specification are defined by the information provided in numeric indicator field <400> followed by the sequence

30 identifier (eg. <400>1, <400>2, etc).

The designation of nucleotide residues referred to herein are those recommended by the IUPAC-IUB Biochemical Nomenclature Commission, wherein A represents Adenine, C represents Cytosine, G represents Guanine, T represents thymine, Y represents a pyrimidine residue, R represents a purine residue, M represents Adenine or Cytosine, K represents Guanine or Thymine, S represents Guanine or Cytosine, W represents Adenine or Thymine, H represents a nucleotide other than Guanine, B represents a nucleotide other than Adenine, V represents a nucleotide other than Thymine, D represents a nucleotide other than Cytosine and N represents any nucleotide residue.

10 Bibliographic details of the publications numerically referred to in this specification are collected at the end of the description.

The flower industry strives to develop new and different varieties of flowering plants.

Altering flower colour has become a particularly important aim in the research and

development undertaken by or on behalf of the flower industry.

Classical breeding techniques have been successfully employed to develop new flowering varieties. However, the generation of plants with desired traits is constrained by, for example, the species' gene pool, a development process which is time consuming and a 20 frequently low success rate. The rapidly increasing sophistication of genetic engineering techniques offers a great opportunity to develop new varieties of plants without some or all of the above constraints.

One important area of flowering plant development is the generation of plants with altered flower colour. Flower colour is predominantly due to two different pigments: flavonoids and carotenoids. The flavonoids contribute to a range of colours from yellow to red to blue. Carotenoids impart an orange or yellow tinge. Particularly important flavonoid molecules include the anthocyanins which are glycosylated derivatives of cyanidin, delphinidin, petunidin, peonidin, malvidin and pelargonidin.

30

The biosynthetic pathway for flavonoid pigments (hereinafter referred to as the "flavonoid

pathway") is now well established (1). The essence of the pathway is a condensation of three molecules of malonyl-CoA with one molecule of *p*-coumaroyl-CoA which is catalysed by chalcone synthase. The product of this reaction, 2', 4, 4', 6'-tetrahydroxychalcone, is generally rapidly isomerised to produce naringenin by chalcone flavanone isomerase. Naringenin is subsequently hydroxylated at the 3' position of the central ring by flavanone 3-hydroxylase to produce dihydrokaempferol (DHK).

The B-ring of the DHK can be hydroxylated at either the 3' or both 3' and 5' positions to produce dihydroquercetin (DHQ) and dihydromyricetin (DHM), respectively. Two key enzymes involved in this pathway are flavonoid 3'-hydroxylase (hereinafter referred to as "F3'H") and flavonoid 3', 5'-hydroxylase (hereinafter referred to as "F3'5'H").

F3'H acts on DHK to produce DHQ and on naringenin to produce eriodicytyol. F3'5'H is a broad spectrum enzyme catalysing hydroxylation of naringenin and DHK in the 3' and 5' positions, in both instances producing pentahydroxyflavanone and DHM, respectively, as well as catalysing hydroxylation of DHQ in the 5' position. The essence of these catalysed reactions is shown in Figure 1 (1).

The ability to control F3'H and F3'5'H activity in plants provides a means to manipulate flower colour. This is successfully described, for example, in International Patent Application Nos. PCT/AU92/00334, PCT/AU93/00127 and PCT/AU94/00265. Levels of F3'H and F3'5'H can be manipulated using genetic means such as altering promoter strength, using anti-sense and ribozyme technologies and through co-suppression. It is important, however, to fully elucidate the endogenous regulatory mechanisms for these and other flavonoid pathway enzymes. Such knowledge can lead to even greater modulation of enzyme activity, especially in plants where expression of, for example, a F3'5'H gene is low.

In work leading up to the present invention, the inventors sought to identify genes involved in the regulation of anthocyanin modification. In accordance with the present invention, the inventors have now identified a molecule which modulates or other

otherwise facilitates activity of a flower cytochrome P450.

The present invention is predicated in part on the isolation of a genetic sequence which encodes a polypeptide which represents a novel class of plant cytochrome b₅ (Cyt b₅) 5 molecules. The genetic sequence of the present invention is referred to and exemplified herein by "difF" which includes a sequence of nucleotides or complementary sequence of nucleotides which encodes the Cyt b₅, i.e., DIF-F, or a mutant, part, fragment, portion thereof or a functional and/or structural equivalent or homologue thereof. For convenience, all such mutants, parts, fragments and portions are referred to herein as a 10 "derivative" or "derivatives". A "derivative" includes mutants, parts, fragments, portions, variants and fusions of the Cyt b₅ protein or corresponding difF gene as well as single or multiple nucleotide substitutions, additions and/or deletions of difF. A "derivative" may also include an agonist or antagonist of Cyt b₅. The term "difF" includes a genomic DNA isolate as well as a cDNA molecule or a chemically prepared molecule generated by the 15 stepwise addition of nucleotides or chemical equivalents thereof. The term "difF" is generically used herein to encompass any molecule encoding a Cyt b₅ or a derivative or homologue thereof and which modulates or otherwise facilitates activity of a Cyt p450 such as but not limited to Cyt P450's involved or otherwise associated with the hydroxylation of a flavonoid compound. An example, the Cyt b₅ or derivative might 20 impact on the activity of a Cyt P450 or may indirectly act via a reductase such as NADPH cytochrome P450 reductase.

Accordingly, one aspect of the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof.

More particularly, the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof wherein said 30 Cyt b₅ modulates or otherwise facilitates activity of a Cyt P450.

Even more particularly, the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof wherein said Cyt b₅ modulates or otherwise facilitates activity of a Cyt P450 wherein the Cyt P450

- 5 comprises the amino acid sequence (F/P/W/L/Y)(G/S/C/N/H/E)X(G/D/A/E)X (R/H/S/K/T) XCX_a(G/A) wherein X is any amino acid and X_a is selected from V, M, A, L, I, P, F or T and wherein the residues in parentheses represent alternatives of a single position.
- 10 Still more particularly, the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof wherein said Cyt b₅ modulates or otherwise facilitates activity of a Cyt P450 enzyme comprising the amino acid sequence (F/P/W/L/Y)(G/S/C/N/H/E)X(G/D/A/E)X
- 15 (R/H/S/K/T) XCX_a(G/A) wherein X is any amino acid and X_a is selected from V, M, A, L, I, P, F or T and wherein the residues in parentheses represent alternatives of a single position and which Cyt P450 is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 20 The preferred Cyt b₅ of the present invention is DIF-F which modulates or otherwise facilitates activity of F3'5'H. However, the present invention extends to other Cyt b₅ molecules which facilitate activity of any Cyt P450 capable of directly or indirectly hydroxylating a flavonoid compound. The present invention is hereinafter described with reference to a Cyt b₅ and its activity on F3'5'H but this is done with the understanding that
- 25 the present invention extends to any member of the Cyt b₅ family especially those involved in the direct or indirect hydroxylation of a flavonoid compound. Examples of other enzymes which are modified or otherwise facilitated by Cyt b₅ include but are not limited to F3'H.
- 30 Although not intending to limit the invention to any one theory or mode of action, Cyt b₅ molecules are believed to interact with cytochrome P450 *via* electrostatic interactions.

These interactions are determined by the primary as well as the tertiary structure of the proteins. An amino acid sequence alignment of petunia Cyt b₅ with other plant Cyt b₅ sequences reveals two regions in the petunia Cyt b₅ sequence that have insertions of six (SELELN) and nine (EDPKPKYLT) amino acids in length. Furthermore, petunia Cyt b₅

- 5 protein has a net positive charge while other plant cytochrome b₅ proteins have a net negative charge. These insertions, which result in a longer petunia Cyt b₅ as well as a change in its charge distribution, may influence the interaction of petunia Cyt b₅ with cytochrome P450 proteins such as petunia F3'5'H.
- 10 The *difF* of the present invention is considered to reside on a separate phylogenetic branch to known Cyt b₅ genes. The preferred novel Cyt b₅ molecule encoded by *difF* comprises the amino acid sequence YKASDDSELELNLVTDSIKEPN or an amino acid sequence having at least about 70% similarity thereto. Even more preferably, the Cyt b₅ of the present invention comprises the amino acid sequence:

15 $[X_1 \ X_2 \X_n] \ KE \ [X_1' \ X_2' \ \ X_{n_1}']F \ [X_1'' \ X_2'' \X''_{n_2}]$ YKASDDSELELNLVTDSIKEPNDSIKEPN $[X_1''' \ X_2''' \X'''_{n_3}] \ EDPKPYLTFVEY$

wherein
$$[X_1 \ X_2 \ ... X_n]$$
, $[X_1' \ X_2' \ ... X_{n_1}']$, $[X_1'' \ X_2'' \ ... X_{n_2}']$ and

- 20 $[X_1^{\prime\prime\prime} X_2^{\prime\prime\prime} X^{\prime\prime\prime}]$ are sequences of any amino acid residues up to n_1 , n_2 and n_3 amino acid residues in length wherein n_1 , $n_2^{\prime\prime}$ and $n_3^{\prime\prime}$ may be the same or different and each is from about 1 to about 200.
- In a preferred aspect of the present invention, there is provided an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof wherein said Cyt b₅ molecule modulates or otherwise facilitates activity of F3'5'H or a derivative, homologue or functional equivalent thereof.
- 30 Preferably, the Cyt b₅ is expressed substantially exclusively in the flower although any Cyt b₅ is contemplated by the present invention provided it modulates or otherwise facilitates

activity of a Cyt P450 molecule. In a particularly preferred embodiment, the Cyt b₅ modulates or otherwise facilitates activity of F3'5'H in flowers.

According to this preferred embodiment, there is provided an isolated nucleic acid

5 molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a Cyt b₅ molecule or a derivative, homologue or functional equivalent thereof wherein said Cyt b₅ modulates or otherwise facilitates activity of F3'5'H or a derivative, homologue or functional equivalent thereof substantially exclusively in flowers.

10 In one particular embodiment, the Cyt b₅ modulates or otherwise facilitates activity of F3'5'H in flowers. In another embodiment, over expression of *difF* may also enhance F3'H activity.

Although not intending to limit the present invention to any one theory or mode of action

15 the Cyt b₅ of the present invention may act at the level of activity of hydroxylating enzyme

(e.g. F3'5'H), at the level of gene transcription (e.g. a transcription regulator) or at the

level of translation. The Cyt b₅ may also act alone or in association with another

molecule. For example, the Cyt b₅ may form a complex with another molecule, e.g. a

reductase, and the Cyt b₅ complex may then act on the hydroxylating enzyme or its genetic

20 sequence. Alternatively, the Cyt b₅ molecule may require the interaction of another

molecule at the level of hydroxylating enzyme or its genetic sequence. In particular, one

of Cyt b₅ or another molecule may interact with the flavonoid hydroxylating enzyme or

genetic sequence encoding same and simultaneously or sequentially, the other of the Cyt b₅

or another molecule may also interact with the flavonoid hydroxylating enzyme or its

25 genetic sequence. The effect(s) of the Cyt b₅ on modulating or otherwise facilitating

activity of the hydroxylating enzyme may require interaction of both Cyt b₅ and the other

molecule.

The preferred Cyt b_5 is DIF-F and comprises the amino acid sequence substantially as set 30 forth in <400>2.

Accordingly, another aspect of the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides or complementary sequence of nucleotides encoding a Cyt b₅ protein or a derivative, homologue or functional equivalent thereof having the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at least about 30% similarity thereto and which Cyt b₅ or derivative, homologue or functional equivalent modulates or otherwise facilitates activity of a flavonoid hydroxylating enzyme such as but not limited to F3'5'H.

The percentage amino acid similarity may be at least about 40%, or at least about 50%, or at least about 60%, or least about 70%, or at least about 80%, or at least about 90-95% or greater to the amino acid sequence set forth in <400>2.

Yet another aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary sequence encoding a Cyt b₅ or a derivative, homologue or functional equivalent thereof wherein the nucleotide sequence is substantially as set forth in <400>1 or a nucleotide sequence having at least 30% similarity thereto or is a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42°C.

- 20 The term "similarity" as used herein includes exact identity between compared sequences at the nucleotide or amino acid level. Where there is non-identity at the nucleotide level, "similarity" includes differences between sequences which result in different amino acids that are nevertheless related to each other at the structural, functional, biochemical and/or conformational levels. Where there is non-identity at the amino acid level, "similarity"
- 25 includes amino acids that are nevertheless related to each other at the structural, functional, biochemical and/or conformational levels. In a particularly preferred embodiment, nucleotide and sequence comparisons are made at the level of identity rather than similarity. Any number of programs are available to compare nucleotide and amino acid sequences. Preferred programs have regard to an appropriate alignment. One such
- 30 program is Gap which considers all possible alignment and gap positions and creates an alignment with the largest number of matched bases and the fewest gaps. Gap uses the

alignment method of Needleman and Wunsch (18). Gap reads a scoring matrix that contains values for every possible GCG symbol match. GAP is available on ANGIS (Australian National Genomic Information Service) at website http://mel1.angis.org.au.

- 5 Preferably, the percentage identity is considered rather than percentage similarity. The term "identity" is used in its broadest sense to include the exact nucleotide or amino acid matches having regard to an appropriate alignment using a standard algorithm. Convenient algorithm in this regard include the Geneworks program (Intelligenetics).
- 10 The percentage nucleotide similarity may be at least about 40%, or at least about 50%, or at least about 60%, or least about 70%, or at least about 80%, or at least about 90-95% or greater to the nucleotide sequence set forth in <400>1.
- Reference herein to a low stringency at 42°C includes and encompasses from at least about 15 0% v/v to at least about 15% v/v formamide and from at least about 1M to at least about 2M salt for hybridisation, and at least about 1M to at least about 2M salt for washing conditions. Alternative stringency conditions may be applied where necessary, such as medium stringency, which includes and encompasses from at least about 16% v/v to at least about 30% v/v formamide and from at least about 0.5M to at least about 0.9M salt for
- 20 hybridisation, and at least about 0.5M to at least about 0.9M salt for washing conditions, or high stringency, which includes and encompasses from at least about 31% v/v to at least about 50% v/v formamide and from at least about 0.01M to at least about 0.15M salt for hybridisation, and at least about 0.01M to at least about 0.15M salt for washing conditions.
- 25 The present invention further provides isolated naturally occurring and recombinant or chemically synthetic forms of DIF-F or other related Cyt b₅ molecules or their derivatives, homologues or functional equivalents thereof. The molecules may be in isolated form or when present in a plant cell. The present invention further extends to antibodies to DIF-F and related Cyt b₅ molecules or their derivatives, homologues or functional equivalents.
- 30 Such antibodies are useful in the immunological detection and/or analysis of plants. The present invention also extends to agonists and antagonists of the Cyt b₅ molecules.

Conveniently, where appropriate, such agonists and antagonists come under the terms "derivative" or "derivatives".

It is proposed in accordance with the present invention that a functional Cyt b₅ (e.g. DIF-F encoded by *difF*) is required for activity of a flavonoid hydroxylating enzyme such as F3'5'H. The term "activity" includes full activity or enhanced, heightened or otherwise facilitated activity. Accordingly, it is further proposed that genetic constructs carrying a nucleotide sequence encoding a flavonoid hydroxylating enzyme such as but not limited to F3'5'H either contain a *difF* or a functional derivative, homologue or equivalent thereof or are used in conjunction with a genetic construct carrying a *difF* or its derivative, homologue or functional equivalent thereof.

Accordingly, another aspect of the present invention contemplates a genetic construct optionally further comprising cistrons encoding one or both of a Cyt P450 or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent of homologue thereof; and/or a reductase or other associated protein.

A single cistron comprises a coding sequence of a particular protein under the control of a promoter sequence. The coding sequence is said to be operably linked to the promoter sequence multiple cistrons may each be under the control of a promoter.

Generally, the Cyt b₅ modulates or otherwise facilitates activity of the Cyt P450 enclosed by the same genetic construct or a Cyt P450 on another genetic construct or encoded by the genome of a host cell.

25

An "associated protein" is a protein which catalyses the transfer of electrons from, for example, a co-enzyme to a prosthetic haem group on reductase. An associated protein may also have a role in facilitating interaction between a Cyt P450 and a reductase.

30 In another embodiment of the present invention, there is provided a genetic construct carrying *difF* or its functional derivative, homologue or equivalent thereof said genetic

construct further comprising a gene for a flavonoid hydroxylating enzyme such as but not limited to a gene encoding a F3'5'H or a functional derivative, homologue or equivalent thereof.

- 5 Another aspect of the present invention provides a transgenic plant or part thereof or cells therefrom comprising genetic material encoding a Cyt b₅ molecule or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof.
- 10 A "part" of a plant includes plant cells and tissues such as petals, flowers, stems, leaves and seeds. Parts of plants include cut or severed flowers.

Yet another aspect of the present invention contemplates a transgenic plant or parts thereof or cells of a transgenic plant, said plant or plant cells comprising genetic material corresponding to *difF* or a functional derivative, homologue or equivalent thereof and optionally a gene encoding an F3'5'H or its derivative, homologue or equivalent.

As stated above, reference to cells of a transgenic plant includes reference to tissues and organs of a plant. Reference to "parts" of a transgenic plant includes flowers (e.g. cut 20 flowers) or flowering plants such as petals.

The present invention also extends to other cells containing or carrying the genetic constructs herein described. Such other cells include yeast cells and bacterial cells.

- 25 In petunia, the alleles encoding F3'5'H are referred to as "hf1" and "hf2". It is proposed, in accordance with the present invention, that difF and/or the product of difF (DIF-F) have a role in facilitating activity of F3'5'H or expression of a gene encoding F3'5'H or its derivatives or homologues or other flavonoid pathway enzymes such as F3'H or its genetic sequences. The present invention extends, however, to the effects of difF or its product
- 30 DIF-F or other related Cyt b₅ molecule on any gene or allele encoding an F3'5'H or a functional derivative, homologue or equivalent thereof. The present invention also

extends, in one particular embodiment, to the effect of difF or DIF-F facilitating or otherwise influencing expression of hf1 and hf2.

Yet another aspect of the present invention provides a method of expressing a nucleotide

5 sequence encoding a Cyt P450 or a functional derivative, homologue or equivalent thereof
in a plant or cells of a plant, said method comprising introducing into said plant or cells of
said plant a genetic construct in single or multicistronic form wherein at least one cistron
encodes a Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or
structural equivalent of homologue thereof; the genetic construct optionally further

10 comprising cistrons encoding one or both of a Cyt P450 or a mutant, part, fragment or
portion thereof or a functional and/or structural equivalent of homologue thereof; and/or a
reductase or other associated protein.

More particularly, the present invention contemplates a method of expressing a genetic sequence encoding an F3'5'H or functional derivative, homologue or equivalent thereof in a plant or cells of a plant, said method comprising introducing a *difF* gene or enhancing expression of a *difF* gene or a functional derivative, homologue or equivalent thereof for a time and under conditions sufficient for the product of *difF* to enhance or otherwise modulate expression of a gene encoding F3'5'H.

20

The present invention further extends to introducing genetic constructs containing separately or together a *difF* and/or a gene encoding F3'5'H or derivatives, homologues or equivalents thereof.

25 Preferably, the modulation of *difF* expression is substantially exclusively in the flowers of plants.

Still another aspect of the present invention provides for the use of a genetic construct comprising a nucleotide sequence encoding a Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof in the manufacture of a plant or cells of a plant in which said Cyt b₅ or a mutant part, fragment

or portion thereof or a functional and/or structural equivalent or homologue thereof enhances, modulates or otherwise facilitates expression of genetic material encoding a Cyt P450 or activity of a Cyt P450.

- 5 Still another aspect of the present invention is directed to the use of *difF* or a functional derivative, homologue or equivalent thereof in the manufacture of a genetic construct capable of enhancing, modulating or otherwise facilitating F3'5'H gene expression or F3'5'H activity.
- 10 The present invention also extends to flowers and cut flowers from transgenic plants and which comprises modified F3'5'H expression levels due to manipulation of *difF* expression. In particular, the present invention is directed to the modulation of flower colour. For example, various shades of blue flowers such as blue roses, carnations and chrysanthemums are contemplated by the present invention.

15

- Although the present invention is particularly directed to the use of *difF* to enhance expression of F3'5'H, the manipulation of certain flower colours may require that the F3'5'H be down regulated. This can be effected by down regulating the expression of *difF* such as by antisense, co-suppression or ribozymes. Alternatively, an endogenous F3'5'H
- 20 may be down regulated by targeting an endogenous *difF* and then an exogenous F3'5'H with an altered substrate specificity or activity introduced to again alter the flow of the metabolites to the flavonoid pathway. All such manipulations and modifications to the methods described therein are contemplated by the present invention.
- 25 The present invention is further described by the following non-limiting Figures and Examples.

In the Figures:

30 **Figures 1a and 1b** are schematic representations of the flavonoid biosynthesis pathways in *P. hybrida* flowers showing the enzymes and genetic loci involved in the conversions.

Enzymes involved in the pathway have been indicated as follows: PAL = phenylalanine ammonia-lyase; C4H = cinnamate 4-hydroxylase; 4CL = 4-coumarate: CoA ligase; CHS= chalcone synthase; CHI= chalcone isomerase; F3H= flavanone 3-hydroxylase; F3'H= flavanoid 3'-hydroxylase; F3'5'H= flavanoid 3'5' hydroxylase; FLS= flavanol synthase;

- 5 DFR= dihydroflavonol-4-reductase; ANS= anthocyanin synthase; 3GT= UDP-glucose: anthocyanin-3-glucoside; 3RT= UDP-rhamnose: anthocyanidin-3-glucoside rhamnosyltransferase; ACT= anthocyanidin-3-rutinoside acyltransferase; 5GT= UDP-glucose: anthocyanin 5- glucosyltransferase; 3' OMT= anthocyanin *O*-methyltransferase; 3', 5' OMT=anthocyanin 3', 5' *O*-methyltransferase. Three flavonoids in the pathway are indicated as: P-3-G= pelargonidin-3-glucoside; DHM=dihydomyricetin;
 - DHQ=dihydroquercetin. The flavonol, myricetin, is only produced at low levels and the anthocyanin, pelargonidin, is rarely produced in *P. hybrida*.
- Figure 2 is a representation showing molecular analysis of difF. (A) Diagram showing the structure of difF. Exons are depicted as thick bars. The triangles indicate the position of dTph1 insertions in the alleles difF-V2082 and difF-W2090. (B) Phylogenetic tree showing the homology of the DIF-F protein to a variety of Cyt b₅ proteins. (C) Alignment of the DIF-F protein with Cyt b₅ from mammals, plants and yeast. Amino acids conserved in more than half of the sequences are indicated by black shading.

20

Figure 3 is a photographic representation showing Northern blot analysis of difF expression. (A) difF expression in different tissues and (B) in the corolla limb at different stages (1-6) of development (2) (C) difF expression in the corolla limb of wild type (R27) and mutant lines (W162, W115, W134) for the regulators an1, an2 and an11. (D) difF mRNA corolla limbs homozygous for the mutable alleles difF-V2082 or difF-W2090 (m/m) and wildtype (+/+) siblings.

Figure 4 is a representation showing analysis of difF mutant flowers. (A) Phenotype of the difF-W2090 allele in a hfl⁺rt⁺ (top) and a hfl⁺rt⁻ background (bottom). (B) PCR 30 analysis of the difF locus in homozygous mutable (m/m) and revertant (+/m) sectors in flowers in different hfl, hf2 and rt genotypes. The intermediately sized fragments are

heteroduplexes that consist of *difF::dTph1* and a *difF*⁺ strand. (C) HPLC analysis of anthocyanin aglycones accumulated in the same sectors. The identity and the molar ratios of the anthocyanin peaks were established by chromatography of pure compounds: del, delphinidin; cya, cyanidin: peo, peonidin; mal, malvidin. (D) F3'5'H enzyme activity in the petal limbs of plants with the indicated phenotype selected from the backcross populations.

Figure 5 is a representation of the nucleotide sequence <400>1 and corresponding amino acid sequence <400>2 of diff. The triangle marks the position of an intron. The underlined sequences mark the insertion sites in the two mutant alleles.

Figure 6a is a diagrammatic representation of the binary plasmid pCGP1280, construction of which is described in Example xx.. Abbreviations are as follows: Tet = the tetracycline resistance gene; LB = left border; RB = right border; surB = the coding region and terminator sequence from the acetolactate synthase gene; 35S = the promoter region from the cauliflower mosaic virus 35S transcript. Restriction enzyme sites are also marked.

Figure 6b to 6g are diagrammatic representations of intermediate plasmids used in the construction of pCGP1280 (Figure 6a). Restriction enzyme sites are also marked. Amp = 20 ampicillin resistance gene.

Figure 6h is a schematic representation of the construction of the binary vector pCGP1280 (Figure 6a). pBS=pBluescript (Stratagene, USA). pANS= antocyanidin synthase promoter, anthocyanidin synthase terminator, Hfl=petunia flavonoid 3'5' hydroxylase cDNA clone, 35S=cauliflower mosaic virus 35S promoter.

Figure 7a is a diagrammatic representation of the binary plasmid pCGP2355, construction of which is described in Example xx.. Abbreviations are as follows: Tet = the tetracycline resistance gene; LB = left border; RB = right border; surB = the coding region and terminator sequence from the acetolactate synthase gene; 35S = the promoter region from the cauliflower mosaic virus 35S transcript. Restriction enzyme sites are also

marked.

Figures 7b to 7d are diagrammatic representations of intermediate plasmids used in the construction of pCGP2355 (Figure 7a). Restriction enzyme sites are also marked. Amp = 5 ampicillin resistance gene.

Figure 7e A schematic representation of the construction of the binary vector pCGP2355 (Figure 7a). AntCHS=Antirrhinum (snapdragon) chalcone synthase promoter difF/DifF=petunia cytochrome b₅ cDNA clone, D8=petunia lipid transfer protein terminator.

Figure 8 is a photograph of transgenic and non-transgenic Exquisite carnation flowers.

Transgenic Exquisite carnations transformed with the T-DNA contained in pCGP 1280 (b and c) produce flower of a similar colour to the non-transformed controls (a and d).

Transgenic Equisite carnations transformed with the T-DNA contained in pCGP2355 (e and f) produce flowers of a novel colour in the violet to deep purple range. Colour photographs are available upon request to the Applicant.

Figure 9 is a representation of an autoradiograph of an RNA blot probed with ³²P-labelled fragments of the *Hfl* cDNA clone contained in pCGP602 (Holton *et al.*, 1993) and diff contained in pCGP2353. A ~1.8 kb Hfl transcript was detected in the petals of the transgenic Exquisite carnations transformed with the T-DNA contained in pCGP2355 (Lanes 2 to 5) or pCGP1280 (Lanes 6 to 9). The same size transcript ws detected in the positive controls of petunia petals of cultivars V30 (Lne 10) and Old Glory Blue (Lane
11). As expected no Hf1 transcript was detected in non-transgenic Exquisite petals (negative control) (Lane 1). A ~0.6 kb difF transcript was only detected in the petals of the transgenic Ezquisite carnations transformed with the T-DNA contained in pCGP2355 (Lanes 2 to 5) and in the positive controls of petunia petals, cultivar V30 and OGB (Lane 10 and 11), respectively. Each lane contained a ~10μg sample of total RNA. A
30 photgraph of the ethidium bromide stained 255 rRNA band is shown as an indication of relative RNA levels.

SUBSTITUTE SHEET (RULE 26)

- 17 -

EXAMPLE 1 GENETIC PROCEDURES

Northern blot, PCR and sequence analyses were done as previously described (2). The petunia lines W138 (relevant genotype: an1-W138, hf1-, hf2-, rt) and V30 (relevant genotype: An1+, Hf1+, Hf2+, Rt+) were maintained as inbred stocks for several generations and were grown under standard greenhouse conditions. Transposon insertion alleles of difF were isolated in the W138 background as previously described (3) using primers complementary to difF and dTphI and maintained by selfing. In the backcrosses of the difF mutant lines with V30, segregation of the unstable anI-W138 and the linked rt allele were scored visually, while the anthocyanin substitution pattern was assayed by TLC and in a few selected plants by HPLC. Segregation of hf1 and hf2 alleles was determined by RFLP analysis (4) and by PCR amplification of the region containing the dTphI insertions for the mutant difF alleles.

15

³²P-Labelling of DNA Probes

DNA fragments (50 to 100 ng) were radioactively labelled with 50 μ Ci of [a-³²P]-dCTP using an oligolabelling kit (Bresatec). Unincorporated [a-³²P]-dCTP was removed by chromatography on a Sephadex G-50 (Fine) column.

20

DNA Sequence Analysis

DNA sequencing was performed using the PRISM™ Ready Reaction Dye Primer Cycle Sequencing Kits from Applied Biosystems. The protocols supplied by the manufacturer were followed. The cycle sequencing reactions were performed using a Perkin Elmer PCR machine (GeneAmp PCR System9600) and run on an automated 373A DNA sequencer (Applied Biosystems).

Homology searches against Genbank, SWISS-PROT and EMBL databases were performed using the FASTA and TFASTA programs (Pearson and Lipman, 1988) or BLAST programs (Altschul *et al.*, 1990). Percentage sequence similarities were obtained using the LFASTA program (Pearson and Lipman, 1988). In all cases ktup values of 6 for nucleotide sequence

comparisons and 2 for amino acid sequence comparisons were used, unless otherwise specified.

Multiple sequence alignments (ktup value of 2) were performed using the ClustalW program incorporated into the MacVectorTM 6.0 application (Oxford Molecular Ltd.).

Low stringency hybridization conditions

Hybridization conditions included a prehybridization step in 10% (v/v) formamide, 1 M NaCl, 10% (w/v) dextran sulphate, 1% (w/v) SDS at 42°C for at least 1 hour. The ³²P-labelled fragments (each at 1x10 cpm/mL) were then added to the hybridization solution and hybridization was continued at 42°C for a further 16 hours. The filters were then washed in 2 x SSC, 1% (w/v) SDS at 42°C for 2 x 1 hour and exposed to Kodak XAR film with an intensifying screen at -70°C for 16 hours.

15 RNA blots

Total RNA was isolated from the petal tissue of Exquisite carnation flowers using an RNAeasy kit from QIAGEN and following the protocols supplied by the manufacturer.

RNA samples were electrophoresed through 2.2 M formaldehyde/1.2% (w/v) agarose gels using running buffer containing 40 mM morpholinopropanesulphonic acid (pH 7.0), 5 mM sodium acetate, 0.1 mM EDTA (pH 8.0). The RNA was transferred to Hybond-N filters (Amersham) as described by the manufacturer.

Hybridization and washing conditions

- 25 RNA blots were probed with ³²P-labelled cDNA fragment (1 x 10⁶ cpm/mL). Prehybridizations (1 hour at 42°C) and hybridizations (16 hours at 42°C) were carried out in 50% (v/v) formamide, 1 M NaCl, 1% (w/v) SDS, 10% (w/v) dextran sulphate. Filters were washed in 2 x SSC, 1% (w/v) SDS at 65°C for 1 to 2 hours and then 0.2 x SSC, 1% (w/v) SDS at 65°C for 0.5 to 1 hour. Filters were exposed to Kodak XAR
- 30 film with an intensifying screen at -70°C for 16 hours.

EXAMPLE 2 CHEMICAL PROCEDURES

Total anthocyanins of flower corolla sectors were extracted and hydrolysed by boiling in 1 ml 2N HCl for 30 min. The anthocyanin aglycones were analysed on a gradient HPLC system equipped with a Vydac C₁₈ reversed phase column (5 μm; 250 x 4.6 mm) and a SPD-M10Avp diode array UV-detector (Shimadzu, Kyoto, Japan). Samples were eluted at 40°C, at a flow rate of 1 ml/min. Anthocyanins were monitored at 547 nm and dihydroflavonols at 280 nm. Solvent system used: a linear gradient elution for 22.5 min from 10 to 75% solvent B (1.5% phosphoric acid, 20% acetic acid and 25% acetonitrile in water) in solvent A (1.5% phosphoric acid in water). Anthocyanins were identified and quantified by comparison with the retention times and peak areas from standards. F3′5′H activity was measured with dihydroquercetin as a substrate as previously described (5), except that formation of the dihydromyricetin product was monitored by HPLC.

EXAMPLE 3 TRANSFORMATION PROCEDURES

20 A. tumefaciens transformations

The plasmids pCGP1280 or pCGP2355 (Figures 6a and 7a) are introduced into the *Agrobacterium tumefaciens* strain AGL0 by adding 5 μg of plasmid DNA to 100 μL of competent AGL0 cells prepared by inoculating a 50 mL MG/L (II) culture and growing for 16 hours with shaking at 28°C. The cells were then pelleted and resuspended in 0.5 mL of 85% v/v 100 mM CaCl₂/15% v/v glycerol. The DNA-*Agrobacterium* mixture was frozen by incubation in liquid N₂ for 2 minutes and then allowed to thaw by incubation at 37°C for 5 minutes. The DNA/bacterial mix was then placed on ice for a further 10 minutes. The cells are then mixed with 1 mL of LB (Sambrook *et al.*, 1989) media and incubated with shaking for 16 hours at 28°C. Cells of *A. tumefaciens* carrying pCGP1280 or pCGP2355 are selected on LB agar plates

containing 50 μ g/mL tetracycline. The presence of pCGP1280 or pCGP2355 is confirmed by Southern analysis of DNA isolated from the tetracycline-resistant transformants.

5 Petunia transformations

(a) Plant Material

Leaf tissue from mature plants is treated in 1.25% w/v sodium hypochlorite for 2 minutes and then rinsed three times in sterile water. The leaf tissue is then cut into 25 mm² squares and precultured on MS media (13) supplemented with 0.05 mg/l 10 kinetin and 1.0 mg/l 2,4-dichlorophenoxyacetic acid (2,4-D) for 24 hours.

(b) Co-cultivation of Agrobacterium Tissue

Agrobacterium tumefaciens strain AGL0 containing genetic material is maintained at 4°C on MG/L agar plates with 100 mg/L gentamycin. A single colony is grown overnight in liquid medium containing 1% w/v Bacto-peptone, 0.5% w/v Bacto-yeast extract and 1% w/v NaCl. A final concentration of 5 x 10⁸ cells/ml is prepared the next day by dilution in liquid MS medium containing B5 vitamins (14) and 3% w/v sucrose (BPM). The leaf discs were dipped for 2 minutes into BPM containing AGL0/genetic material. The leaf discs are then blotted dry and placed on co-cultivation media for 4 days. The co-cultivation medium consists of SH medium (15) supplemented with 0.05 mg/l kinetin and 1.0 mg/l 2,4-D and included a feeder layer of tobacco cell suspension spread over the co-cultivation medium with a filter paper placed on top of the tobacco cell suspension.

25 (c) Recovery of transgenic plants

After co-cultivation, the leaf discs are transferred to a selection medium (MS medium supplemented with 3% w/v sucrose, α-benzylaminopurine (BAP) 2 mg/l, 0.5 mg/l α-naphthalene acetic acid (NAA), kanamycin 300 mg/l, 350 mg/L cefotaxime and 0.3% w/v Gelrite Gellan Gum (Schweizerhall)). Regenerating explants are transferred to fresh selection medium after 4 weeks. Adventitious shoots which survive the kanamycin selection are isolated and transferred to BPM containing 100

mg/l kanamycin and 200 mg/l cefotaxime for root induction. All cultures are maintained under a 16 hour photoperiod (60 μ mol. m⁻², s⁻¹ cool white fluorescent light) at 23 ± 2°C. When roots reach 2-3 cm in length the transgenic petunia plantlets are transferred to autoclaved Debco 51410/2 potting mix in 8 cm tubes.

5 After 4 weeks, plants are replanted into 15 cm pots, using the same potting mix, and maintained at 23 °C under a 14 hour photoperiod (300 μ mol. m⁻², s⁻¹ mercury halide light).

Flower Colour coding

10 The colour codes are taken from the Royal Horticultural Society's Colour Chart (RHSCC).

EXAMPLE 4

TRANSFORMATION OF DIANTHUS CARYOPHYLLUS CV. EXQUISITE

The binary vectors pCGP1280 or pCGP2355 were introduced into *A. tumefaciens* strain AGL0 cells, as described in Example . The pCGP1280/AGL0 or pCGP2355/AGL0 cells were used to transform carnation plants.

20 (a) Plant Material

15

Dianthus caryophyllus (cv. Exquisite) cuttings are obtained Baguely F & I, Flower and Plant Growers, Heatherton Road, Clayton South, Victoria, Australia. The outer leaves are removed and the cuttings are sterilised briefly in 70% v/v ethanol followed by 1.25% w/v sodium hypochlorite (with Tween 20) for 6 min and rinsed three times with sterile 25 water. All the visible leaves and axillary buds are removed under the dissecting microscope before co-cultivation.

(b) Co-cultivation of Agrobacterium and Dianthus Tissue

Agrobacterium tumefaciens strain AGL0 (19), containing the binary vector pCGP1280 30 or pCGP2355, is maintained at 4°C on LB agar plates with 50 mg/L tetracycline. A single colony is grown overnight in liquid LB broth containing 50 mg/L tetracycline and

diluted to 5 x 10⁸ cells/mL next day before inoculation. *Dianthus* stem tissue is co-cultivated with *Agrobacterium* for 5 days on MS medium supplemented with 3% w/v sucrose, 0.5 mg/L BAP, 0.5 mg/L 2,4-dichlorophenoxy-acetic acid (2,4-D), 100 mM acetosyringone and 0.25% w/v Gelrite (pH 5.7).

5

(c) Recovery of Transgenic Dianthus Plants

For selection of transformed stem tissue, the top 6-8 mm of each co-cultivated stem is cut into 3-4 mm segments, which were then transferred to MS medium (13) supplemented with 0.3% w/v sucrose, 0.5 mg/L BAP, 0.5 mg/L 2,4-D, 1 μg/L 10 chlorsulfuron, 500 mg/L ticarcillin and 0.25% w/v Gelrite. After 2 weeks, explants are transferred to fresh MS medium containing 3% w/v sucrose, 0.16 mg/L thidiazuron (TDZ), 0.5 mg/L indole-3-butyric acid (IBA), 2 μg/L chlorsulfuron, 500 mg/L ticarcillin and 0.25% w/v Gelrite and care is taken at this stage to remove axillary shoots from stem explants. After 3 weeks, healthy adventitious shoots are transferred to hormone free MS medium containing 3% w/v sucrose, 5 μg/L chlorsulfuron, 500 mg/L ticarcillin, 0.25% w/v Gelrite. Shoots which survive 5 μg/L chlorsulfuron are transferred to the same medium for shoot elongation.

Elongated shoots are transferred to hormone-free MS medium containing 5 μg/L 20 chlorsulfuron, 500 mg/L ticarcillin and 0.4% w/v Gelrite, in glass jars, for normalisation and root production. All cultures are maintained under a 16 hour photoperiod (120 mE/m²/s cool white fluorescent light) at 23± 2°C. Normalised plantlets, approximately 1.5-2 cm tall, were transferred to soil (75% perlite/25% peat) for acclimation at 23°C under a 14 hour photoperiod (200 mE/m²/s mercury halide light) for 3-4 weeks. Plants 25 were fertilised with a carnation mix containing 1g/L CaNO₃ and 0.75 g/L of a mixture of microelements plus N:P:K in the ratio 4.7:3.5: 29.2.

EXAMPLE 5

TRANSFORMATION OF ROSA HYBRIDA

1. Rosa hybrida

5 Plant tissues of the rose are transformed according to the method disclosed in PCT/AU91/04412, having publication number WO92/00371.

2. Rosa hybrida

a. Plant Material

10 Kardinal shoots are used. Leaves are removed and the remaining shoots (5-6 cm) are sterilized in 1.25 % w/v sodium hypochlorite (with Tween 20) for 5 minutes followed by three rinses with sterile water. Isolated shoot tips are soaked in sterile water for 1 hour and precultured for 2 days on MS medium containing 3% w/v sucrose, 0.1 mg/L BAP, 0.1 mg/l kinetin, 0.2 mg/l Gibberellic acid, 0.5% w/v polyvinyl 15 pyrrolidone and 0.25% w/v Gelrite Gellan Gum, before co-cultivation.

b. Co-cultivation of Agrobacterium and Rosa shoot Tissue

Agrobacterium tumefaciens strains ICMP 8317 (18) and AGL0, containing genetic a particular construct are maintained at 4°C on MG/L agar plates with 100 mg/L

- 20 gentamycin. A single colony from each Agrobacterium strain is grown overnight in liquid MG/L broth. A final concentration of 5 x 10^8 cells/ml is prepared the next day by dilution in liquid MG/L. Before inoculation, the two Agrobacterium cultures are mixed in a ratio of 10:1. A longitudinal cut is made through the shoot tip and an aliquot of 2 μ l of the mixed Agrobacterium cultures is placed as a drop on the shoot
- 25 tip. The shoot tips are co-cultivated for 5 days on the same medium used for preculture.

Agrobacterium tumefaciens strain AGL0 is maintained at 4°C on MG/L agar plates with 100 mg/L kanamycin. A single colony from each Agrobacterium strain is grown overnight in liquid MG/L broth. A final concentration of 5 x 10⁸ cells/ml is prepared the next day by dilution in liquid MG/L.

c. Recovery of Transgenic Rosa Plants

After co-cultivation, the shoot tips are transferred to selection medium. Shoot tips are transferred to fresh selection medium every 3-4 weeks. Galls observed on the shoot tips are excised when they reached 6-8 mm in diameter. Isolated galls are transferred 5 to MS medium containing 3% w/v sucrose, 25 mg/l kanamycin, 250 mg/l cefotaxime and 0.25% w/v Gelrite Gellan Gum for shoot formation. Shoots regenerated from gall tissue are isolated and transferred to selection medium. GUS histochemical assay and callus assay are used to identify transgenic shoots. Transgenic shoots are transferred to MS medium containing 3% w/v sucrose, 200 mg/l cefotaxime and 10 0.25% w/v Gelrite Gellan Gum for root induction. All cultures are maintained under 16 hour photoperiod (60 μE cool white fluorescent light) at 23 \pm 2°C. When the root system is well developed and the shoot reached 5-7 cm in length the transgenic rose plants are transferred to autoclaved Debco 514110/2 potting mix in 8 cm tubes. After 2-3 weeks plants are replanted into 15 cm pots using the same potting mix and 15 maintained at 23 °C under a 14 hour photoperiod (300 μE mercury halide light). After 1-2 weeks potted plants are moved to glasshouse (Day/Night temperature : 25-28°C/14°C) and grown to flowering.

EXAMPLE 6

20 TRANSFORMATION OF CHRYSANTHEMUM MORIFOLIUM

a. Plant Material

Chrysanthemum morifolium cuttings are obtained. Leaves are removed from the cuttings, which were then sterilized briefly in 70% v/v ethanol followed by 1.25% 25 w/v sodium hypochlorite (with Tween 20) for 3 minutes and rinsed three times with sterile water. Internodal stem sections are used for co-cultivation.

b. Co-cultivation of Agrobacterium and Chrysanthemum Tissue

Agrobacterium tumefaciens strain LBA4404 (Hoekema et al 1983), containing is 30 grown on MG/l agar plates containing 50 mg/l rifampicin and 10 mg/l gentamycin. A single colony from each Agrobacterium is grown overnight in the same liquid

medium. These liquid cultures are made 10% v/v with glycerol and 1 ml aliquots transferred to the freezer (-80°C). A 100-200µl aliquot of each frozen Agrobacterium is grown overnight in liquid MG/l containing 50 mg/l rifampicin and 10 mg/l gentamycin. A final concentration of 5 x 10⁸ cells/ml is prepared the next day by 5 dilution in liquid MS containing 3% w/v sucrose. Stem sections are co-cultivated with Agrobacterium in co-cultivation medium for 4 days.

c. Recovery of Transgenic Chrysanthemum Plants

After co-cultivation, the stem sections were transferred to selection medium. After 3-4 weeks, regenerating explants are transferred to fresh medium. Adventitious shoots which survive the kanamycin selection are isolated and transferred to MS medium containing kanamycin and cefotaxime for shoot elongation and root induction. All cultures are maintained under a 16 hour photoperiod (80 μE cool white fluorescent light) at 23 ± 2°C. Leaf samples are collected from plants which rooted on kanamycin and Southern blot analysis is used to identify transgenic plants. When transgenic chrysanthemum plants reach 4-5 cm in length, they are transferred to autoclaved Debco 51410/2 potting mix in 8 cm tubes. After 2 weeks, plants are replanted into 15 cm pots using the same potting mix and maintained at 23°C under a 14 hour photoperiod (300 μE mercury halide light). After 2 weeks potted plants are 20 moved to glasshouse (Day/Night temperature : 25-28°C/14°C) and grown to flowering.

EXAMPLE 7 PLANTS

25

The Exquisite carnation was employed for transformation experiments (Baguley F&I, Flower & Plant Growers, Heatherton Road, Clayton South, Victoria, Australia). this carnation has a bicoloured flower.

EXAMPLE 8 IDENTIFICATION OF difF

To identify additional genes involved in anthocyanin modification, the inventors isolated cDNAs corresponding to genes that are down-regulated in flowers with a mutation in the regulatory anthocyanin-1 (an1) gene (6). Based on its sequence, one gene, potentially involved in flavonoid hydroxylation, was chosen for detailed analysis. This gene, termed herein "difF", encodes a polypeptide of 149 amino acids that represents a novel class of plant Cyt b₅ proteins (Fig. 2b, c). The highest degree of similarity is clustered around the pairs of histidine residues (His-39 and His-63) that correspond to the axial ligands for heme binding (7). Although the Cyt b₅ sequences show strong divergence in the C-terminal part of the polypeptide, they have a strikingly similar hydropathy plot. This hydrophobic C-terminal part anchors the enzyme to the endoplasmic reticulum (ER) membrane (7).

15

EXAMPLE 9 EXPRESSION OF difF

To examine the function of *difF* in anthocyanin biosynthesis, its expression pattern 20 was analysed by Northern blots and compared to the expression pattern of the *dfr* gene, encoding dihydroflavonol 4-reductase, a key enzyme of the anthocyanin pathway. Fig. 3A shows that the *difF* transcripts accumulate in the limb and tube of the flower corolla and in the ovaries, but not in vegetative organs such as leaves, root and stem. During petal development the temporal *difF* expression pattern closely matches that of the gene encoding dihydroflavonal reductase *(dfr)*, with both transcripts reaching a maximum around stage 3, when the flower bud starts to open (Fig. 3B). To test if *difF* expression is controlled by any of the known regulators of the anthocyanin pathway, the inventors analysed *difF* transcript levels in stage 3 flowers (2) of the corresponding mutants. Fig. 3C shows that *difF* expression is 30 down-regulated in petal limbs of *an1*, *an2* and *an11* mutants, when compared to wildtype. Although *an2-W115* is a null allele, this mutation reduces anthocyanin

synthesis strongly, but does not block it completely. This indicates that an2 function is partially redundant and explains the residual difF and dfr transcripts detected in an2-W115 petal limbs (Fig. 3C). Taken together, these data show that the spatio-temporal and genetic control of difF expression are consistent with a role in 5 athocyanin synthesis.

EXAMPLE 10 ISOLATION OF diff MUTANTS

10 To establish the in vivo function of difF, the inventors isolated difF mutants by a PCR based screen (3) to identify plants of the line W138 in which a dTph1 transposon had inserted in the difF gene. Among 4000 W138 plants, the inventors found that two individuals that were heterozygous for the wildtype $difF^+$ allele and a transposon insertion derivative (difF-V2082 and difF-W2090 respectively). 15 germinated from these individuals, that had been produced by self-pollination, and progeny homozygous for difF-V2082 and difF-W2090 identified by PCR. Sequence analysis showed that in difF-V2082 a 284 bp dTph1 element had inserted in the first exon, 10 bp upstream of the splice-site, thereby disrupting the protein coding sequence. The difF-W2090 allele contained a 284 bp dTph1 insertion in the middle 20 of exon 2, that also disrupts the coding sequence, see Figure 5 for mapping of Northern analysis showed that flowers of difF-V2082 homozygous insertions. progeny accumulated difF transcripts that are about 300 bp larger than the wildtype difF transcript (Fig. 3D). By analogy to other dTph1 insertion alleles this mutant transcript is likely to contain the transcribed dTph1 sequence. In difF-W2090 25 homozygotes the amount of difF mRNA was reduced about three-fold when compared to difF+ siblings. Since difF-W2090 is relatively unstable, these transcripts most likely result from dTph1 excisions and probably contain different transposon footprints.

EXAMPLE 11 MUTATIONS OF diff AFFECTS FLOWER COLOUR

To study the effects of mutant difF alleles into an $hf1^+$ or $hf2^+$ genetic background, 5 the inventors made backcrosses with line V30 ($hf1^+$, $hf2^+$, $an1^+$, rt^+), using difFmutant lines as the recurrent parent. As expected, these progenies (co-)segregated 1:1 for $an1^{mutable}$ mutable (anT) and rt plants (Table 1). If the 5' substitution of anthocyanin is dependent on the segregation of hf1 and hf2 alone, one would expect to find plants accumulating malvidin ($hf1^+ hf2^+$ and hf^+ , $hf2^-$), malvidin plus peonidin 10 (hf1, hf2+; the relatively weak hf2 locus enables the 5' substitution of only about 50% of the anthocyanins) or peonidin (hfl- hf2-) corolla pigments in a ratio 2:1:1. However, combined results of the two backcross populations segregating for difF-V2082 and difF-W2090, respectively showed a segregation ratio of 38:51:6 (Table 1). This suggested that a third mutant gene segregated that reduced the 5' substitution, 15 possibly diff. To test this directly, the inventors subjected representative plants of the various phenotypic classes to Southern blot and PCR analyses to determine the hf1, hf2 and difF genotype. This revealed that the malvidin accumulating plants were all hfl+ difF+, while those accumulating a mixture of malvidin and peonidin were either $hf1^+$, $difF^m$, $hf1^-hf2^+$ $difF^m$ or $hf1^-hf2^+difF^+$.

20

Closer inspection showed that the hfl^+ rt^+ individuals which were homozygous for the difF-W2090 allele had variegated flowers with purple (revertant) sectors and spots on a purplish magenta (mutant) background (Fig. 4A top). Also flowers of $hfl^ hf2^+$ $difF^m$ siblings were variegated, although the colour difference between mutant and revertant tissue was less pronounced. In an hfl^+ rt plants the variegation was seen as "dull-grey" revertant spots and sectors on a "dull-red" mutant background (Fig. 4A bottom). To test whether these variegated flower colours were due to genetic instability of the difF-W2090 allele, the inventors isolated DNA from several large revertant petal sectors and from the mutant corolla sectors and analysed the difF gene 30 by PCR. Fig. 4B shows that reversion of the flower colour are associated with (somatic) excision of the dTphl element from difF-W2090. Also difF-V2068

5

individuals had variegated flowers, but the frequency of revertant spots was lower by at least one order of magnitude.

EXAMPLE 12

MUTATION OF diff REDUCES MODIFICATION OF THE ANTHOCYANIN

To examine how the difF mutation affected flower colour, the inventors dissected (isogenic) difF⁺ revertant and difF mutant sectors of single flowers and analysed the 10 anthocyanin aglycones by HPLC. Some representative chromatograms are shown in Fig. 4C. In $difF^+$ revertant petal sectors on hf1 rt plants about 80% of the anthocyanins are 3', 5' substituted (delphinidin), while in difm mutant sectors of the same flower this amount is reduced to about 40% (Fig. 4C, top). The reduced delphinidin accumulation is correlated with an increase in the accumulation of 3' 15 substituted and anthocyanin (cyanidin) from 20 to 63%. This indicates that the difF mutation reduced the formation of 3', 5' hydroxylated anthocyanins by about 50% and that the remaining precursors are converted into a 3' hydroxylated anthocyanin. The same phenomenon was observed in hfl+, rt+ flowers. In this background, the difF-W2090 mutation reduced the fraction of 3', 5' substituted anthocyanins (malvidin) 20 from 94 to 73%, which correlated with an increase in 3' substituted anthocyanin (peonidin) from 6 to 27% (Fig. 4C, middle). This indicated a 25% inhibition in the formation of 3', 5' substituted anthocyanins. In $hf1^-hf2^+$ tissue less than half of the anthocyanins were 3',5' substituted (44%), possibly because the hf2 locus expresses lower amounts of F3'5'H protein, or a F3'5'H protein with lower activity. In this 25 background a diff mutation decreased 3',5' substitution further down to 29%, corresponding to about 35% inhibition (Fig. 4C, bottom).

EXAMPLE 13 diff MUTATION REDUCES F3'5'H ACTIVITY

30

To test if difF stimulates 3', 5' substitution of anthocyanin precursors by regulating

the activity of the CytP450 enzyme F3'5'H, the inventors measured F3'5'H activities in different genotypes. Because this required larger quantities of petal tissue, these measurements could not be performed on (isogenic) mutant and revertant sectors of single flowers. Instead, the inventors selected two or three plants from the V30 backcross populations for each genotype and determined F3'5'H enzyme activity in microsomes that were isolated from stage 4 petal limbs. Fig. 4D shows that the *difF* mutation reduced *hf1* encoded F3'5'H activity by about 20-fold, while *hf2* encoded F3'5'H activity was reduced approximately 3-fold.

- 10 The data show unequivocally that *in vivo*, a Cyt b₅ (DIF-F) is required for activity of Cyt P450, F3′5′H, without an apparent effect on other Cyt P450 enzymes, such as those involved in 3′-hydroxylation of dihydroflavonols, synthesis of the flavonoid precursor cinnamic acid, or synthesis of hormones controlling plant development. Both *in vitro* reconstruction experiments (7) as well as *in vivo* over-expression experiments in yeast (8) and human cells (9) have shown that the activity of a Cyt P450 can be increased 10 to 20 fold by co-expression of a Cyt b₅. Therefore, *difF* may provide a critical tool to increase the activity of a *f3′5* ħ transgene in ornamental flowers that normally lack blue colours.
- 20 Those skilled in the art will appreciate that the invention described herein is susceptible to variations and modifications other than those specifically described. It is to be understood that the invention includes all such variations and modifications. The invention also includes all of the steps, features, compositions and compounds referred to or indicated in this specification, individually or collectively, and any and 25 all combinations of any two or more of said steps or features.

EXAMPLE 14

SCREENING FOR EFFECTS OF CYT B₅ ON P450 MOLECULES

The effects of Cyt b₅ molecules on different P450 molecules such as F3'5'H or F3'H 5 is determined using yeast. Details of suitable yeast strains and expression vectors is sshown in US Patent No. 5,349,125. In one embodiment, the petunia Cyt b₅ is incorporated into the genome of a yeast and genetic constructs encoding a Cyt P450 introduced into the cell. Expression of Cyt P450 may be measured by any number of ways. In relation to F3'5'H, for example, radiolabelled dihydrokoenpferol (DHK) or radiolabelled naringinin may be used. For other P450's, the product or substrate may be measured using, for example, HPLC, TLC or other suitable procedures.

Exquisite is a carnation cultivar that produces bi-coloured flowers with a deep red centre and a pale pink rim. The petals normally accumulate cyanidin, a 3',4'-15 hydroxylated anthocyanidin and the flavonolos quercetin and kaempferol (a 3',4'-hydroxylated flavonol and 4'-hydroxylated flavonol, respectively).

Introduction of a petunia flavonoid 3'5' hydroxylase (F3'5'H) under the control of a carnation ANS (anthocyanidin synthase) promoter (contained in pCGP1280 [Figure 20 6a]) results in either no or a slight alteration of petal colour with low levels of delphinidin (3',4',5'-hydroxylated anthocyanidin) [Figure 1a and 1b] (Tables 2 and 3) being produced.

Introduction of the same chimeric F3'5'H (ANS:HF1:ANS) along with diff under the control of a snapdragon CHS (chalcone synthase) promoter both contained in pCGP2355 (Figure 7a) resulted in a major shift in flower colour. The flowers of the transgenic Exquisite/pCGP2355 flowers were deep purple with a pale purple rim (figure 8). HPLC analysis of the anthocyanidins and flavonol content of Exquisite/pCGP2355 showed that delphinidin (the 3',4',5'-hydroxylated anthocyanidin) was the predominant anthocyanidin produced.

This result suggested that expression of the introduced difF along with the F3'5'H chimeric gene enhanced F3'5'H activity so that higher levels of delphinidin were produced compared to the expression of the F3'5'H chimeric gene with the absence of $cytb_5$.

5

SUMMARY W138(diff) X V30 CROSSES

	Table 1: Number of plants with flower coloured and anthogyanths in the Parakiyases intociding N v301X N138 white with coloured spots (anl [®]) .:difF®	f plants with flower colour and white with coloured spots (anl ^m)	ed spol	s (anl ^m)	- IIOCYAIII	114 811	אפרטרוב הפרטרוב	full co	loured	full coloured (An1 [±]) ::diff	A MI38
JBS	Rr+			_tt			Br +			_ #	
UIII CEOSS	mal mal/peo	o peo	del	Jel del/cya	cva	ma l	mal/peo peo del del/cya	oad	del	del/cya	Cya
H 22363		44			\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	<	4	6		^ :-
SHEET	3	0	15	14	, ,	16	22	, m	0		^ O
18) T	· · · · · · · · · · · · ·	r	6	! ! ! ! ! !	\	· · · >	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	61	1	8 8 8 8 8	^!!
JLE 26	<3 1	\	19	.9 12	17	12	5929	۸ ۳	0	1 0	^ O
total	>	121		1	^ / !	· · · · · · · · · · · · · · · · · · ·)11	01	-	^ }
	4	<	34	34 36	24	38	38 51	9	/ 0	2 0	^ d
*backcros.	*backcross family 22363 segregated for diff-V2082, and 22364 segregated for diff-W2090	segregated	for d	ifF-V2082,	and 223(54 segre	egated for	di FF-W20	060		

~
<u>ده</u>
\equiv
್ಷ
_

1		_				Т			7							Τ	1	Т	7					ļ	1	Т	\neg
Σ	%	0.0%	0.0%	0.0%		3.8%		4.1%	0.0%	5.9%	4.7%	4.2%	11.5%	3.8%		ì	1.9%	14.8%	4.6%	%0:0	6.5%	4.9%	4.7%	4 7%	è	4.2%	4.7%
쏘	mg/g	1.09	2.20	0.95		2.16		1.45	2.25	1.50	1.59	1.58	1.66	1.72			2.97	0.92	1.14	1.04	0.97	0.81	96.0	0 80	30.0	C/.D	1.06
0	mg/g	68.0	1.28	0.65		1.22		0.72	0.92	0.89	92.0	1.07	09.0	0.93			1.71	0.56	0.83	0.63	0.87	0.63	0.95	0.67	20.0	0.68	98.0
М	3/3w	0.00	0.00	0.00		0.13		0.09	0.00	0.14	0.11	0.11	0.26	0.10			0.37	0.22	0.09	0.00	0.12	0.07	0.09	0.07	70.0	0.00	0.09
Del	%	%0.0	0.0%	%0.0		18.4%		58.2%	%6.0	88.8%	71.1%	67.4%	%0.68	63.8%			89.3%	80.2%	52.1%	0.7%	70.3%	60.4%	54.0%	70 00	40.770	53.9%	27.6%
Cya	mg/g	1.23	1.03	0.63	20:0	1.13		0.27	1.69	80.0	0.18	0.19	0.08	0.36			0.30	0.26	0.61	1.39	0.34	0.35	0.40	5	0.03	0.36	0.59
Del	mg/g	0.00	00.0	8	3	0.25		0.38	0.02	0.65	0.45	0.38	0.64	0.63	3		2.47	1.04	99.0	0.01	0.81	0.54	0.47		0.43	0.42	0.80
RHSCC		56a		560	204												71a	71a	71a	187c	83a	79a	703	377	64a	79a	71a
Acc#		20173 (h)	20173 (h)	20172 (4)	(0) 5/107	20183	6107	19787	19788	19789	19794	19796	19809	10812	17017		19810	19810	19812	19788	19789	19794	10706	17/70	19802	19805	19806
Construct	Consulation		connoi	control	control		Ans-HII-Ans	A == 11f1 A == / Ant CHS_cuth 5-D8	Aus-Hill-Aus/Aut/HS-cyth5-D8	Ans-Hft Ans/Ant/HS-cyth5-D8	And Hell Ans/Ant/HS-cyth5-D8	Alls-Hit-Alls/Allectic Cycle 20	Ans-rill-Ans/Anctio-cyco Co	Ans-Hil-Aus/Aut.Cits-cyto-Do	Ans-Hil-Ans/AntCH3-cyu3-Do		Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hfl-Ans/AntCHS-cytb5-D8	And Hell Ans/Ant/HS_cyth5_D8	Ans-Mis-mis/micros of control of the Mis-Mis-Mis-Mis-Mis-Mis-Mis-Mis-Mis-Mis-	Aus-Hill-Austricing June 1	Alls-full-Australian-cycle Do	Ans-Ht1-Ans/AntCH3-cyt03-Do	2355 inner Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hfl-Ans/AntCHS-cytb5-D8	
a557	pcdr		ınner	mir	whole		1280 rım		IIIII CC67	2355 mm	11111 CCC7	mn ccs2	Z355 mm	2355 rim	2355 rim		2355 inner	2355 inner	2255 inner	23.00	2325 Inner	2325 Inner	7322 inner	2355 inner	2355 inner	2355 inner	2355 inner
	Cultivar		Exquisite	Exquisite	Exquisite		Exquisite		Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite		Evanieite	Daminite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Evanicite	Exquisite

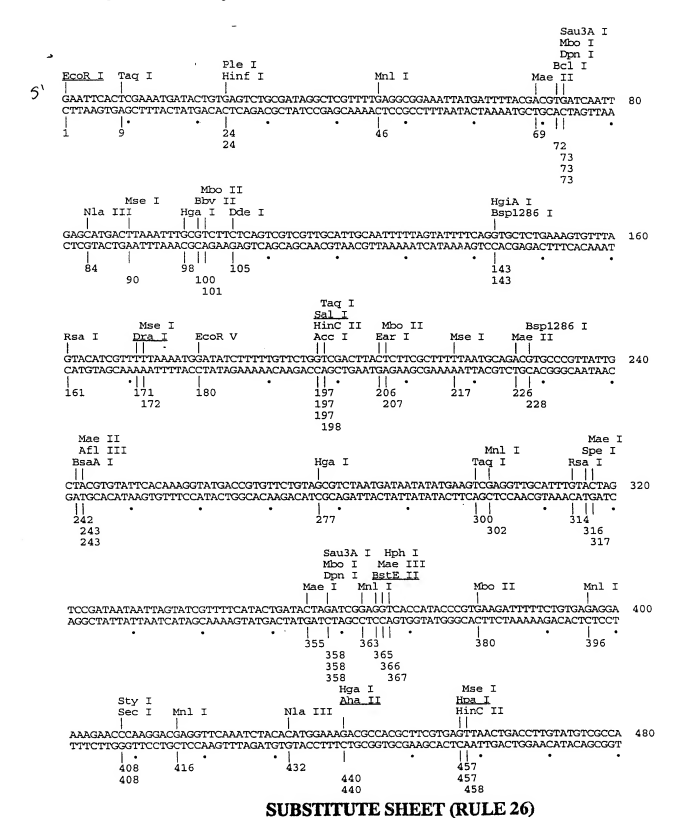
Fable 3

	_	_	_	_	_	_																	
	X	%	%0.0	%0 0	%0 0	%0 0	%0.0	2.8%	0.0%	%0.0	%0 0	0.00	5 70%	769/	5 00 S	5 20%	702.9	2007	6.20%	11 3%	%0.0	0.0%	0.0%
ω.	Ж	mg/g	1.71	1.65	2.06	1.13	1.78	2.50	2.00	2.02	1.98	1 20	1 36	1 45	211	1 07	2.48	3 05	200	0 37	2.38	1.75	1.86
Flavonols	~	mg/g	0.88	0.89	2.24	0.86	1.17	1.36	98.0	1.32	96.0	0.72	0.85	0.95	=	= ==	1.57	1 64	1 15	0.42	2.52	1.27	1.48
	M	g/gm	0.00	0.00	0.00	0.00	0.00	0.11	0.00	0.00	0.00	0.19	0.12	0.11	0.16	0.17	0.27	0.28	0.20	0.09	0.00	0.00	0.00
lins	Del	%	0.8%	0.7%	0.7%	4.1%	1.3%	24.8%	8.5%	1.7%	%9.0	86.9%	63.0%	67.4%	52.5%	63.5%	74.7%	89.9%	75.6%	71.9%	%0.0	%0.0	%0.0
Anthocyanidins	Cya	mg/g	1.10	1.36	1.40	0.87	1.23	1.27	1.75	1.60	1.28	0.12	0.42	0.27	99.0	0.56	0.62	0.19	0.41	0.25	1.68	1.40	1.44
Ar	Del	mg/g	0.01	0.01	0.01	0.04	0.02	0.42	0.16	0.03	0.01	0.81	0.71	0.56	0.73	0.97	1.83	1.73	1.26	0.64	0.00	0.00	0.00
	RHSCC		64p	64a	64a	74a	64a	61a	61a	64a	61a	79a	72a	71a	64a	71a	71a	71a	71a	71a	56a	56a	56a
	Acc#		20186	20196	20181	20184	20194	20198	20189	20194	19786	19818	19815	19804	19802	19806	19808	19810	19808	19819	21129	21129	21129
	Construct		Ans-Hfl-Ans	Ans-Hf1-Ans	Ans-Hf1-Ans	Ans-Hf1-Ans	Ans-Hf1-Ans	Ans-Hf1-Ans	Ans-Hfl-Ans	Ans-Hf1-Ans	Ans-Hfl-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hfl-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	Ans-Hfl-Ans/AntCHS-cytb5-D8	Ans-Hfl-Ans/AntCHS-cytb5-D8	Ans-Hf1-Ans/AntCHS-cytb5-D8	control	control	control
	bCGP		1280	1280	1280	1280	1280	1280	1280	1280	2355	2355	2355	2355	2355	2355	2355	2355	2355	2355	Aglo	Aglo	Aglo
	Cultivar		Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite	Exquisite								

TABLE 4....

ANSpromoter -> Restriction Map

DNA 5 quence 2552 b.p. GAATTCACTCGA ... TCATAATCTAGA linear



871 871 871

ANSpromoter -> Restriction Map Nla III HinP I Hha I NspH I Dde I Hae II Nsp7524 I Hph I PflM I Fok I Afl III Esp I Eco47 III || • 547 520 530 492 509 483 547 484 492 547 493 493 Fnu4H I Bbv I Alu I <u>Pvu II</u> Ple I SfaN I Mae I Taq I Xca I <u>Gsu</u> I Hinf I Hph I Mae II NspB Acc I HIIGTTGTTCTTACATTTAGGTGAAAGACGTTTCTCCAGCTGCTAGGAGTCGAGATGCGAAATTGTCGTTTGCGACTGTATAC CAACAAGAATGTAAATCCACTTTCTGCAAAGAGGTCGACGATCCTCAGCTCTACGCTTTAACAGCAAACGCTGACATATG | | •| 604 111 635 577 **5**85 594 600 607 635 604 595 596 596 Nla III Sph I NspH I Nsp7524 I Nsi I SfaN I BsmA I Tth111 I Fok I Mbo II 111 11 CTTGATAGAAATĠĠÁTĠĆATGCAAGTAAAGAAGGTAŤCTTCTAATTCATCTTTCGTAĠAGACATAGCGTGAATTTGĠACG GAACTATCTTTACCTACGTACGTTCATTTCTTCCATAGAAGATTAAGTAGAAAGCATCTCTGTATCGCACTTAAACCTGC ||| || • 653 698 717 654 655 657 657 657 658 Mae II <u>SnaB I</u> BsaA I Alu I Hph I GGGTCTTTGGTTTGAGAAAGATAACAGCTTTACGTATTTTTGTAGATGGGTGAAACCTTTTCAAATCCGTATAAGCGTAA 800 CCCAGAAACCAAACTCTTTCTATTGTCGAAATGCATAAAAACATCTACCCACTTTGGAAAAGTTTAGGCATATTCGCATT | 746 769 751 - 751 752 Sau3A I Mbo I Dpn I SfaN I BsmA I Bcl I Bsr I AGACGACAACTGGGCTTTAGGGGACACATTCTTTCAGGTATAATTGATGCGACTAACAATAGTCTCCACTGATCATATTC TCTGCTGTTGACCCGAAATCCCCTGTGTAAGAAAGTCCATATTAACTACGCTGATTGTTATCAGAGGTGACTAGTATAAG 846 862 870 809

- 38 -ANSpromoter -> Restriction Map Mbo II Taq I Ear I Mae II Acc I Mae II TACTCTTCTACGTTCGATACTGACTGTTTCTGGTTATTTGGTAGACAGGAGATTATTTGGACGTAGCAATTCAGTAGCGT 960 ATGAGAAGATGCAAGCTATGACTGACAAAGACCAATAAACCATCTGTCCTCTAATAAACCTGCATCGTTAAGTCATCGCA 883 890 921 941 884 894 Mae II Afl III Mae I <u>Pml I</u> Sty I BsaA I Sec I Afl III Avr II AGAGATGTTTCCACACGTGTTATCGTAAAAGAAGCAAGATAAGCCTAATGCCTAGGGTGGTGGTATGACTTCCGTTGCTT 1040 TCTCTACAAAGGTGTGCACAATAGCATTTTCTTCGTTCTATTCGGATTACGGATCCCACCACCATACTGAAGGCAACGAA 1011 974 1011 974 1011 975 1012 975 Sau3A I Mbo I Dpn I <u>Pvu I</u> Taq I Mbo II Nla III Cla I Tth111 II Mse I Mae I BSpH I ATCGATCGTGCTTGTAAGTAATTTCCGTCTTATCTTTTCCTGTTATATAAAGTTAATCTTCTCTAGGACTTTCATGAACC 1120 TAGCTAGCACGAACATTCATTAAAAGGCAGAATAGAAAAGGACAATATTTTCAATTAGAAGAAGATCCTGAAAGTTACTTTCG . 1041 1049 1093 i103 1112 1042 1097 1113 1043 1044 1044 1044 Mae I Sau3A I Spe I Mbo I Xca I Dpn I Alu I Acc I Nla III Mae I Mse I Mae II Taq I TIGTTTGTGTATTTATTTCTCGATCAACATGATAGAGCTAGTTTTTAAGCAACGTATACTAGTAGTCTATTGGAAGTTAA 1200 AACAAACACATAAATAAAGAGCTAGTTGTACTATCTCGATCAAAAATTCGTTGCATATGATCATCAGATAACCTTCAATT 1 | • 1148 1158 1165 1122 1140 1172 1142 1156 1174 1142 1174 1142 1178 1179 Sau3A I Mbo I Nla III Dpn I NspH I Alw I Mae I Ple I Mse I Rsa I Nsp7524 I Mbo II Hinf I 11 GACACGGTTCŤTAAAAAGGTACGATCCAAGTGAAGCATGTTAGATATGACACTTŤCTTCTAGGGACGACTCTCGTATGCC 1280 $\tt CTGTGCCAAGAATTTTTCCATGCTAGGTTCACTTCGTACAATCTATACTGTGAAAGAAGATCCCTGCTGAGAGGCATACGG$ 1235 | |• 1211 1219 1255 1267 1223 1235 1259 1267 1223 1236 1223 1223 Fok I SfaN I BsmA T Nsi I ACCCGACTTTTCAATTTTTTTTGTGAATGTTAGATGTGTGTATATAATGCATCCGAAAGATGTCTCAACGAACAAATGA 1360 TGGGCTGAAAAAGTTAAAAAAAACACTTACAATCTACACACATATATTACGTAGGCTTTCTACAGAGTTGCTTGTTTACT 1 11

SUBSTITUTE SHEET (RULE 26)

1328

1330

1343

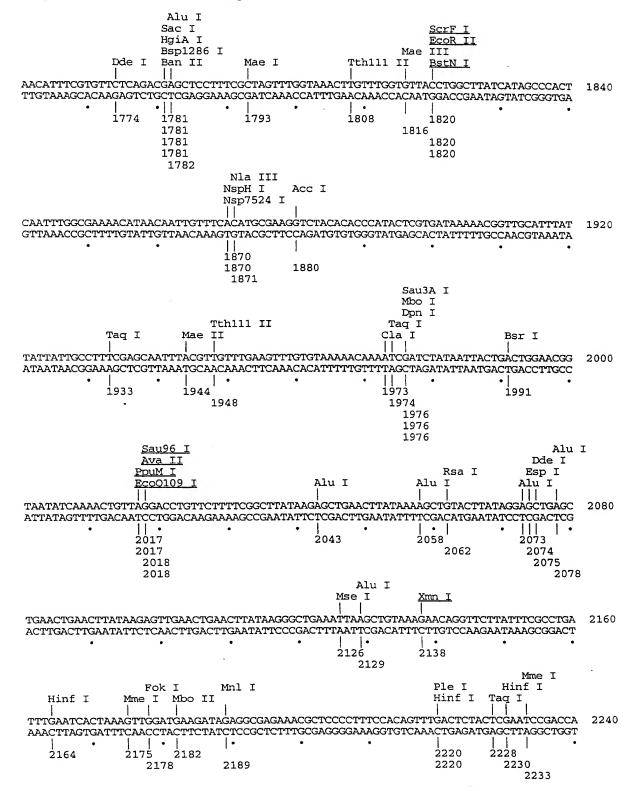
ANSpromoter -> Restriction Map Sau3A I Sau3A I Mbo I Mbo I Dpn I Mse I Dpn I Bcl I Mse I Ase I Taq I GCCACCTACTTCGATCACTCGCTATCAATGTTATTAATGCCTTGTTGATTTAATAGTTGATCAATAATAGTAAAAATCTA 1440 $\tt CGGTGGATGAAGCTAGTGAGCGATAGTTACAATAATTACGGAACAACTAAAATTATCAACTAGTTATTATCATTTTAGAT$ || 1393 • | | 1371 1411 1419 1420 1373 1394 1420 1373 1373 Alu I Sac I HgiA I Bsp1286 I Nla IV Mae I Ase I Ban I Taq I Dde I Mse I Ban II Mse I Mae III Hga I BsmA I 11 TTCAAGGGTATAGTCTCCCGTTCACACTCATCGGGGTTACACTAGCGAGCTCCATTAATCGGTGCCTTAATCGAGACGCT 1520 AAGTTCCCATATCAGAGGGCAAGTGTGAGTAGCCCCAATGTGATCGCTCGAGGTAATTAGCCACGGAATTAGCTCTGCGA || • 11 | | | | | | | | | | | | • , 1515 1511 'i495 1487 1476 1507 1453 1501 1519 1482 1494 1487 1501 1513 1487 1487 1488 Nla III Sty I Sec I Sau3A I Nco I Mbo I Dsa I Mae I HinP I Tthlll I Hha I Dpn I Hae III Alw I Taq I Mae I <u>Hae I</u> Hae II Nla III AAGAACTATACCATGACCTAGTCAGCGCCATGGGACTGATGTAGGCCACACAATCTCGATGATCCGAAAACGCTAGAGTT TTCTTGATATGGTACTGGATCAGTCGCGGTACCCTGACTACATCCGGTGTGTTAGAGCTACTAGGCTTTTGCGATCTCAA 1544 • 1563 **1576** 1532 1581 1564 1535 1545 1538 1581 1545 1581 1548 1581 1548 1548 1548 1549 Sty I Sec I Nco I Mae I Dsa I BsmA I EcoR V Tag I Nla III Taq I Drd I Mae I BsmA I Mae III BstU I 1607 1614 11. 1664 1623 1637 1666 1619 1640 1612 1670 1618 1618 1618 1618 Mbo II Mae I Bbv II Nde I Mse I Rsa I Alu I AATAAGTCTTGTGTACGATGGGTAGCTAGTGAATTAAAGGTAATCACTTTACTCGTGTTCACAAGAAGACCATTCATATG TTATTCAGAACACATGCTACCCATCGATCACTTAATTTCCATTAGTGAAATGAGCACAAGTGTTCTTCTGGTAAGTATAC | | 1704 1745 1755 1714 1693

SUBSTITUTE SHEET (RULE 26)

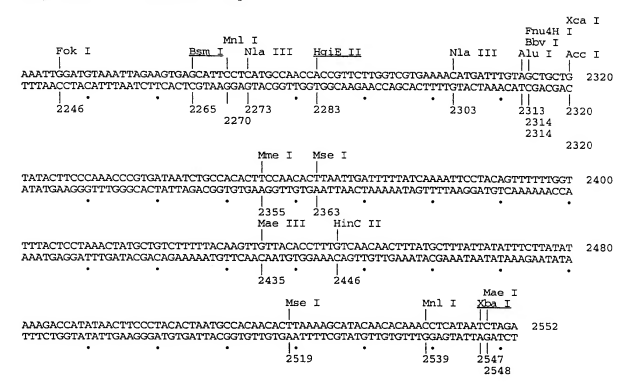
1706

1745

ANSpromoter -> Restriction Map



ANSpromoter -> Restriction Map



BIBLIOGRAPHY

- 1. Holten, T. Drug Metabolism and drug interactions 12: 359-368, 1995.
- 2. de Vetten, N. et al. Genes Dev. 11: 1422-1434, 1997.
- 3. Koes, R. et al. Proc. Natl. Acad. Sci. USA 81: 8149-8153, 1995.
- 4. Holton, T. A. et al. Nature 336: 276-279, 1993.
- 5. Menting et al. Plant Physiol. 106: 633-642, 1994.
- 6. Kroon et al. Plant J. 5: 69-80, 1994.
- 7. Vergeres, G & Waskell, L. Biochimie 77: 604-620, 1995.
- 8. Auchus et al. J. Biol. Chem. 273: 3158-3165, 1998.
- 9. Aoyama et al. Proc. Natl. Acad. Sci. USA 87: 5425-5429, 1990.
- 10. G. Vergeres & L. Waskell. Biochimie 77: 604-620, 1995.
- 11. Ito et al. Journal of Bacteriology 153: 163-168, 1983.
- 12. Gerfinkel and Nester. Journal of Bacteriology 144: 732-743, 1980.
- 13. Sambrook *et al.* (eds) *Molecular Cloning: A Laboratory Manual* (2nd ed). Cold Spring Harbor Laboratory Press, USA, 1989.
- 14. Murashige and Skoog. Plant Physiology 15: 73-97, 1962.
- 15. Gamborg et al. Experimental Cell Research 50: 151-158, 1968.
- 16. Schenk and Hilderbrandt. Canadian Journal of Botany 50: 199-204, 1972.
- 17. Jefferson et al. EMBO Journal 6(13): 3901-3907, 1987.
- 18. Needleman and Wunsch, *J. Mol. Biol.* 48: 443-453, 1970

CLAIMS:

- 1. An isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a cytochrome b_5 (Cyt b_5) molecule or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof.
- 2. An isolated nucleic acid molecule according to claim 1 wherein the Cyt b₅ modulates or otherwise facilitates activity of a cytochrome P450 molecule (Cyt P450).
- 4. An isolated nucleic acid molecule according to claim 3 wherein the Cyt P450 molecule is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 5. An isolated nucleic acid molecule according to claim 4 wherein the Cyt P450 is flavonoid 3',5'-hydroxylase (F3',5'H).
- 6. An isolated nucleic acid molecule according to claim 4 wherein the Cyt P450 is flavonoid 3'-hydroxylase (F3'H).
- 7. An isolated nucleic acid molecule according to any one of claims 1 to 6 wherein the Cyt b_5 comprises the amino acid sequence:

YKASDDSELELNLVTDSIKEPN

SUBSTITUTE SHEET (RULE 26)

or an amino acid sequence having at least 70% similarity thereto.

8. An isolated nucleic acid molecule according to claim 7 wherein Cyt b₅ comprises the amino acid sequence:

$$[X_1X_2 ... X_n]KE[X_1', X_2' ... X_{n1}']$$

 $F[X_1'', X_2'' ... X_{n2}'']$

YKASDDSELELNLVTDSIKEPNDSIK

EPN[X₁"', X₂"' ... X"'_{n3}] EDPKPYLTFVES

wherein $[X_1, X_2 ... X_n]$, $[X_1', X_2' ... X'_{n1}]$, $[X''_1, X_2'' ... X''_{n2}]$ and $[X_1''', X_2''' ... X'_{n2}]$ are amino acid sequences of any amino acid residues up to n, n_1 , n_2 and n_3 amino acid residues in length wherein n, n_1 , n_2 and n_3 may be the same or different and each is from about 1 to about 200.

- 9. An isolated nucleic acid molecule according to any one of claims 1 to 8 wherein the Cyt b_5 comprises the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at least about 30% similarity thereto.
- 10. An isolated nucleic acid molecule according to any one of claims 1 to 9comprising the nucleotide sequence substantially as set forth in <400>1, or a nucleotide sequence having at least about 30% similarity thereto or a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42° C.
- 11. A genetic construct in single or multicistronic form wherein at least one cistron encodes a Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or structural equivalent of homologue thereof; the genetic construct optionally further comprising cistrons encoding one or both of a Cyt P450 or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent of homologue thereof; and/or a reductase or other associated protein.
- 12. A genetic construct according to claim 11 wherein the Cyt b₅ modulates or

otherwise facilitates activity of the Cyt P450 encoded by the same genetic construct or a Cyt P450 on another genetic construct or encoded by the genome of a host cell.

- 14. A genetic construct according to claim 13 wherein the Cyt P450 molecule is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 15. A genetic construct according to claim 14 wherein the Cyt P450 molecule is F3',5'H.
- 16. A genetic construct according to claim 14 wherein the Cyt P450 molecule is F3'H.
- 17. A genetic construct according to any one of claims 11 to 16 wherein the Cyt b₅ comprises the amino acid sequence:

Y K A S D D S E L E L N L V T D S I K E P N or an amino acid sequence having at least 70% similarity thereto.

18. A genetic construct according to claim 17 wherein the Cyt b₅ comprises the amino acid sequence:

$$[X_1X_2 ... X_n]KE[X_1', X_2' ... X_{n1}']$$

 $F[X_1'', X_2'' ... X_{n2}'']$

SUBSTITUTE SHEET (RULE 26)

YKASDDSELELNLVTDSIKEPNDSIK EPN[X₁"', X₂"' ... X"'_{n3}] EDPKPYLTFVES

wherein $[X_1, X_2 ... X_n]$, $[X_1', X_2' ... X'_{n1}]$, $[X''_1, X_2'' ... X''_{n2}]$ and $[X_1''', X_2''' ... X''_{n2}]$ are amino acid sequences of any amino acid residues up to n, n_1 , n_2 and n_3 amino acid residues in length wherein n, n_1 , n_2 and n_3 may be the same or different and each is from about 1 to about 200.

- 19. A genetic construct according to any oen of claims 11 to 18 claim 17 or 18 wherein the Cyt b_5 comprises the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at least about 30% similarity thereto.
- 20. A genetic construct according to any one of claims 11 to 19 wherein the Cyt b_5 is encoded by a nucleotide sequence substantially as set forth in <400>1, or a nucleotide sequence having at least about 30% similarity thereto or a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42° C.
- 21. A transgenic plant or part thereof or cells therefrom comprising genetic material encoding a Cyt b₅ molecule or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof.
- 22. A transgenic plant or part thereof or cells therefrom according to claim 21 wherein the Cyt b_5 modulates or otherwise facilitates activity of a Cyt P450.
- 23. A transgenic plant or part thereof or cells therefrom according to claim 21 or 22 wherein the Cyt P450 molecule comprises the amino acid sequence $(F/P/W/L/Y)(G/S/C/N/H/E)X(G/D/A/E)X(R/H/S/K/T) XCX_a(G/A)$ wherein X is any amino acid and X_a is selected from V, M, A, L, I, P, F or T and wherein the residues in parentheses represent alternatives of a single position.

- 24. A transgenic plant or part thereof or cells therefrom according to claim 23 wherein the Cyt P450 molecule is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 25. A transgenic plant or part thereof or cells therefrom according to claim 24 wherein the Cyt P450 is F3',5'H.
- 26. A transgenic plant or part thereof or cells therefrom according to claim 24 wherein the Cyt P450 is F3'H.
- 27. A transgenic plant or part thereof or cells therefrom according to any one of claims 21 to 26 wherein the Cyt b₅ comprises the amino acid sequence:

Y K A S D D S E L E L N L V T D S I K E P N or an amino acid sequence having at least 70% similarity thereto.

28. A transgenic plant or part thereof or cells therefrom according to claim 27 wherein Cyt b₅ comprises the amino acid sequence:

$$[X_1X_2 ... X_n]KE[X_1',X_2' ... X_{n1}']$$

 $F[X_1'', X_2'' ... X_{n2}'']$

YKASDDSELELNLVTDSIKEPNDSIK

wherein $[X_1, X_2 ... X_n]$, $[X_1', X_2' ... X_{n1}']$, $[X_1', X_2'' ... X_{n2}']$ and $[X_1'', X_2''' ... X_{n2}'']$ are amino acid sequences of any amino acid residues up to n, n_1 , n_2 and n_3 amino acid residues in length wherein n, n_1 , n_2 and n_3 may be the same or different and each is from about 1 to about 200.

29. A transgenic plant or part thereof or cells therefrom according to claims 21 to 28 wherein the Cyt b_5 comprises the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at latest about 30% similarity thereto.

- 30. A transgenic plant or part thereof or cells therefrom according to any one of claims 21 to 29 or 28 or 29 comprising the nucleotide sequence substantially as set forth in <400>1, or a nucleotide sequence having at least about 30% similarity thereto or a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42° C.
- 31. A method of expressing a nucleotide sequence encoding a Cyt P450 or a functional derivative, homologue or equivalent thereof in a plant or cells of a plant, said method comprising introducing into said plant or cells of said plant a genetic construct in single or multicistronic form wherein at least one cistron encodes a Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or structural equivalent of homologue thereof; the genetic construct optionally further comprising cistrons encoding one or both of a Cyt P450 or a mutant, part, fragment or portion thereof or a functional and/or structural equivalent of homologue thereof; and/or a reductase or other associated protein.
- 32. A method according to claim 31 wherein the Cyt b₅ modulates or otherwise facilitates activity of the Cyt P450 encoded by the same genetic construct or a Cyt P450 on another genetic construct or encoded by the genome of a host cell.
- 33. A method according to claim 31 or 32 wherein the Cyt P450 molecule comprises the amino acid sequence (F/P/W/L/Y)(G/S/C/N/H/E)X(G/D/A/E)X $(R/H/S/K/T) XCX_a(G/A)$ wherein X is any amino acid and X_a is selected from V, M, A, L, I, P, F or T and wherein the residues in parentheses represent alternatives of a single position.
- 34. A method according to claim 33 wherein the Cyt P450 molecule is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 35. A method according to claim 34 wherein the Cyt P450 molecule is

F3',5'H.

- 36. A method according to claim 34 where the Cyt P450 molecule is F3'H.
- 37. A method according to any one of claims 31 to 36 wherein the Cyt b_5 comprises the amino acid sequence:

Y K A S D D S E L E L N L V T D S I K E P N or an amino acid sequence having at least 70% similarity thereto.

38. A method according to claim 37 where the Cyt b₅ comprises the amino acid sequence:

$$[X_1X_2 ... X_n]KE[X_1', X_2' ... X_{n1}']$$

 $F[X_1'', X_2'' ... X_{n2}'']$

YKASDDSELELNLVTDSIKEPNDSIK EPN[X,"', X,"' ... X"',] EDPKPYLTFVES

wherein $[X_1, X_2 \dots X_n]$, $[X_1', X_2' \dots X_{n1}']$, $[X_1', X_2'' \dots X_{n2}'']$ and $[X_1''', X_2''' \dots X_{n2}'']$ are amino acid sequences of any amino acid residues up to n, n_1 , n_2 and n_3 amino acid residues in length wherein n, n_1 , n_2 and n_3 may be the same or different

39. A method according to any one of claims 31 to 38 where the Cyt b_5 comprises the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at latest about 30% similarity thereto.

and each is from about 1 to about 200.

- 40. A method according to any one of claims 31 to 39 where the Cyt b_5 is encoded by a nucleotide sequence substantially as set forth in <400>1, or a nucleotide sequence having at least about 30% similarity thereto or a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42° C.
- 41. Flowers cut or severed from a plant according to any one of claims 21 to

30.

- 42. Reproductive parts of a plant according to any one of claims 21 to 30.
- 43. Use of a genetic construct comprising a nucleotide sequence encoding a Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof in the manufacture of a plant or cells of a plant in which said Cyt b₅ or a mutant part, fragment or portion thereof or a functional and/or structural equivalent or homologue thereof enhances, modulates or otherwise facilitates expression of genetic material encoding a Cyt P450 or activity of a Cyt P450.
- 44. Use according to claim 43 wherein the Cyt P450 molecule comprises the amino acid sequence (F/P/W/L/Y)(G/S/C/N/H/E)X(G/D/A/E)X(R/H/S/K/T) $XCX_a(G/A)$ wherein X is any amino acid and X_a is selected from V, M, A, L, I, P, F or T and wherein the residues in parentheses represent alternatives of a single position.
- 45. Use according to claim 45 wherein the Cyt P450 molecule is involved in or otherwise associated with the direct or indirect hydroxylation of a flavonoid compound.
- 46. Use according to claim 46 wherein the Cyt P450 is flavonoid 3',5'-hydroxylase F3',5'H.
- 47. Use according to claim 46 wherein the Cyt P450 is F3'H.
- 48. Use according to any one of claims 44 to 47 wherein the Cyt b_5 comprises the amino acid sequence:

Y K A S D D S E L E L N L V T D S I K E P N or an amino acid sequence having at least 70% similarity thereto.

49. Use according to claim 49 wherein Cyt b₅ comprises the amino acid sequence:

$$[X_1X_2 \dots X_n]KE[X_1', X_2' \dots X_{n1}']$$

 $F[X_1", X_2" \dots X_{n2}"]$

YKASDDSELELNLVTDSIKEPNDSIK

 ${\rm E\;P\;N\;[X_1"',\;X_2"'\;...\;X"'_{n3}]\;E\;D\;P\;K\;P\;Y\;L\;T\;F\;V\;E\;S}$

wherein $[X_1, X_2 ... X_n]$, $[X_1', X_2' ... X'_{n1}]$, $[X''_1, X_2'' ... X''_{n2}]$ and $[X_1''', X_2''' ... X''_{n2}]$ are amino acid sequences of any amino acid residues up to n, n_1 , n_2 and n_3 amino acid residues in length wherein n, n_1 , n_2 and n_3 may be the same or different and each is from about 1 to about 200.

- 50. Use according to any one of claims 43 to go wherein the Cyt b_5 comprises the amino acid sequence substantially as set forth in <400>2 or an amino acid sequence having at latest about 30% similarity thereto.
- 51. Use according to any one of claims 43 to 50 comprising the nucleotide sequence substantially as set forth in <400>1, or a nucleotide sequence having at least about 30% similarity thereto or a nucleotide sequence capable of hybridizing to <400>1 under low stringency conditions at 42° C.

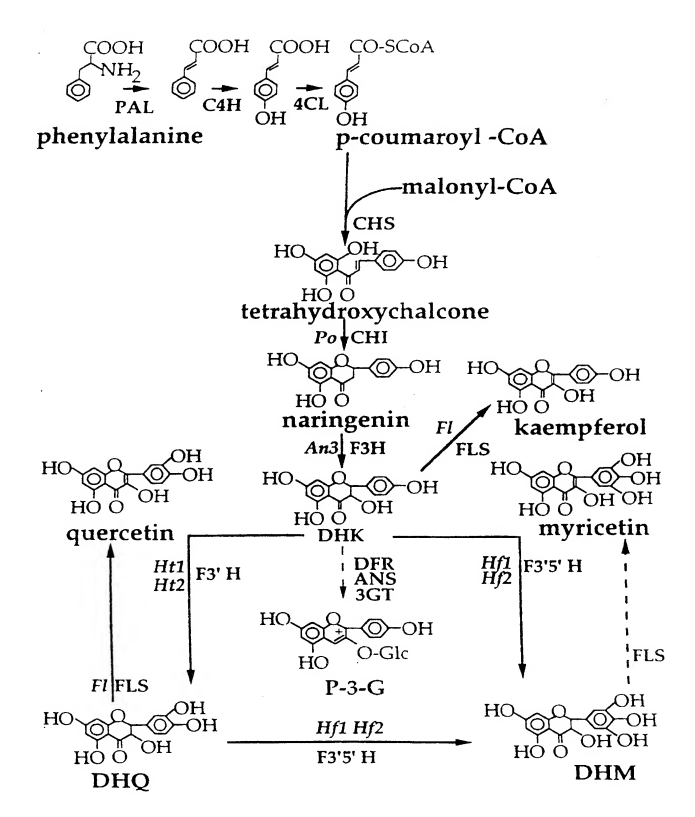
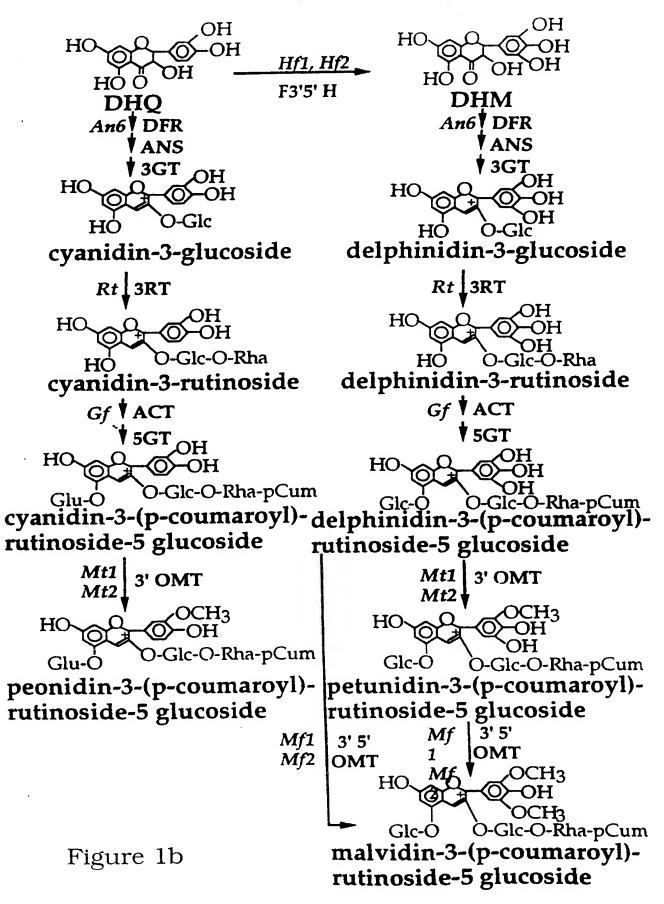


Figure 1a
SUBSTITUTE SHEET (RULE 26)



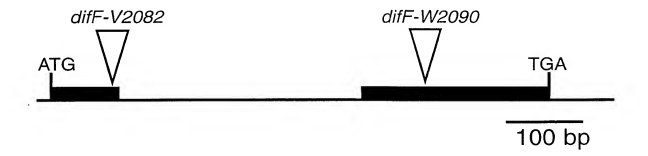


Figure 2a

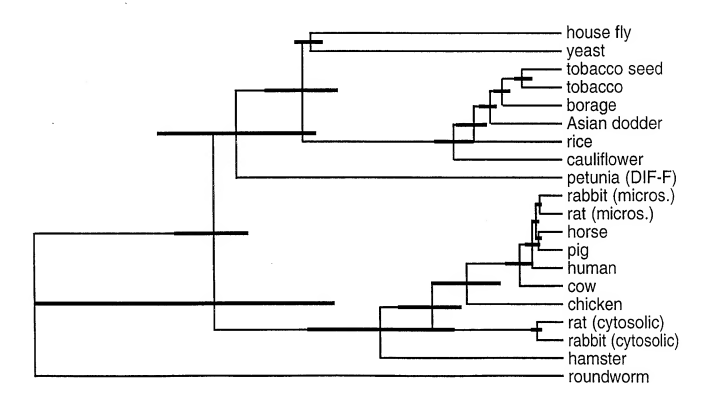


Figure 2b **SUBSTITUTE SHEET (RULE 26)**

335 335 339 33	80 75 73	07 15 12 00	4 8 8 8 8 9 8 9 9 9 9 9 9 9 9 9 9 9 9 9
-50000-50000000-50000-6000000000-0		-0	-0
	ススのエエー でえたよほ	Z Z I	
XXXXO	H H X H U	8 1 H > 1	
HH>>>	X Z L O X	01>>1	
	ស ស ដ ដ ដ	HIXEI	
>	дддΣн	HIKII	
>>>>>	BBZKK	HIXII	1 1 14 4 1
* * K K K K	K K K K H	нижии	N I O N Z
H H Z S Q	**************************************	1 1 7 8 8 8	ωι « Οι
нанны	HHXXD	1 1 4 1 5	EH I U X I
ннн>н	တတ္ကတ္လ	1124	\Rightarrow \vdash \vdash \vdash \vdash
аанан		11200	中またこれ
3333	00000	11400	东道东二镇
E E O O E	>>H>H		> 1 22 22 1
SOOS		11221	нижии
S M S M G		വാനമാനമ	4 1 H H 1
ZHWZU	ZZEIOS	ыырон	וטאול
ZZZZZ	医思比口尼	SEHZS	
	HHHHH		או אמא
对对因公司	REERE	医肾正己胺	4 1 4 2 2
OMANA		E H Z S H	о I & O O
H H >>>	00000 00XXO	K K E K S	K I LE H I
ы ы о « О	ಷಕ್ಕ∺೮	SSHFD	العابد
дыныя	DONAH	阴阴冒火民	и п ц н и
HHHHO	医原原の口	$\sigma \sigma \omega \mapsto \omega$	агына
东西置东京	K K H H Z	00000	1 1 2 4 4
** >	дддан	I I S T X P P D T T	H T C I I
>>	NAME E	1 1 4 4 1	116161
AUOHA	医医蛋白口	11411	1160>
米末杯 日 ι	(11 (2) (2) (2)	1 1 > 1 1	ここはより
8 D D D I	# # # # # © © © © ©	ומטוו	HTK1 N
σαισι	0,0,0,0,0	ннооо.	ZIMMH
M OI I Z I	8	HHHH H >	27 24 1
HE I HI			33 1 04 1 1
T T T T T T T T T T T T T T T T T T T	11111X 21212	ннн>н	S W W T N W
cow rabbit petunia tobacco yeast	cow rabbit petunia tobacco yeast	cow rabbit petunia tobacco yeast	cow rabbit petunia tobacco yeast
cow rabbit petunia tobaccc yeast	f obi cun oac ast	cow rabbit petunia tobacco	w bbi tur bac
cow raby pet top	cow rabl peti tob	cow rabk petu toba	cow rabb petu toba

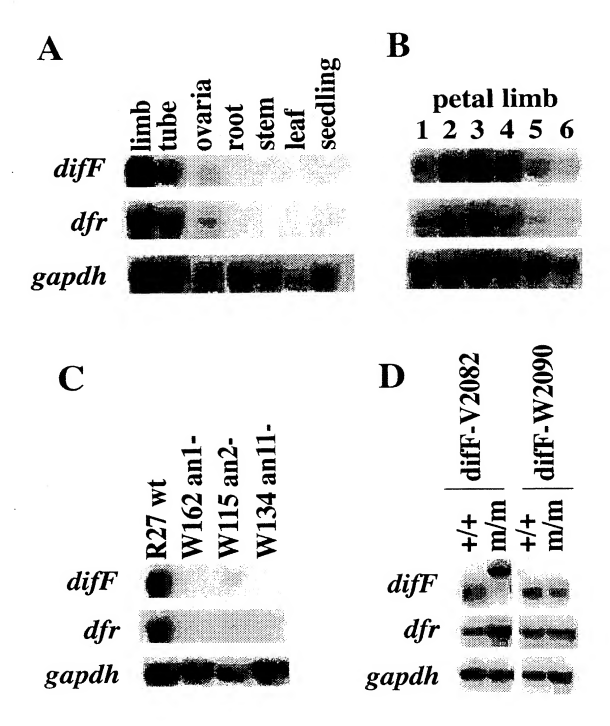
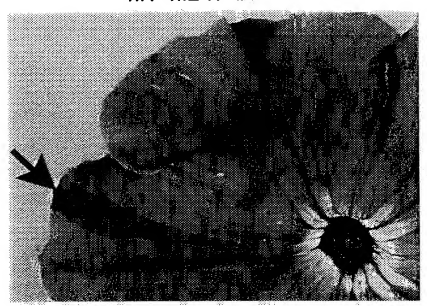


Figure 3
SUBSTITUTE SHEET (RULE 26)

hf1+hf2 rt+difFm



hf1+hf2 rf difFm



Figure 4a
SUBSTITUTE SHEET (RULE 26)

$$\frac{hf1^{+}hf2^{-}rf}{m/m} + \frac{hf1^{+}hf2^{-}rf^{+}}{m/m} + \frac{hf1^{-}hf2^{+}rf^{+}}{m/m} + \frac{hf1^{-}hf2^{+}rf^{+}}{m/m} + \frac{difF::dTph1}{-difF^{+}}$$

Figure 4b

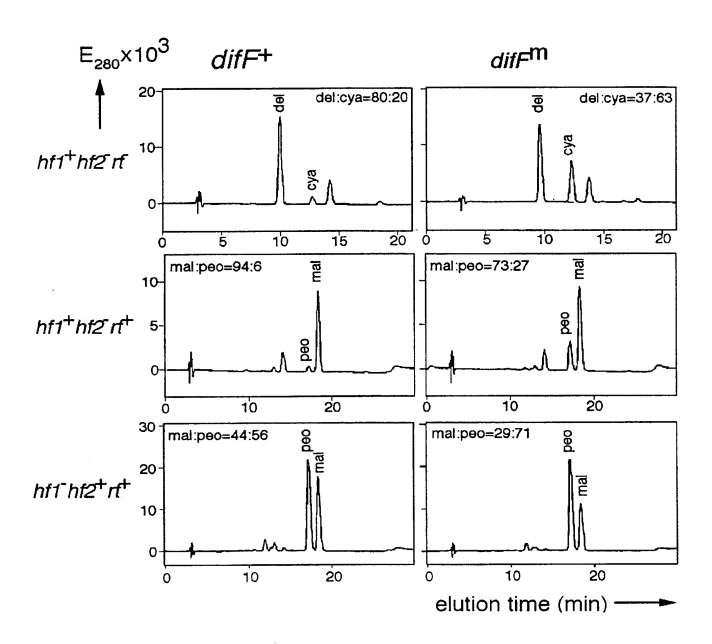


Figure 4c

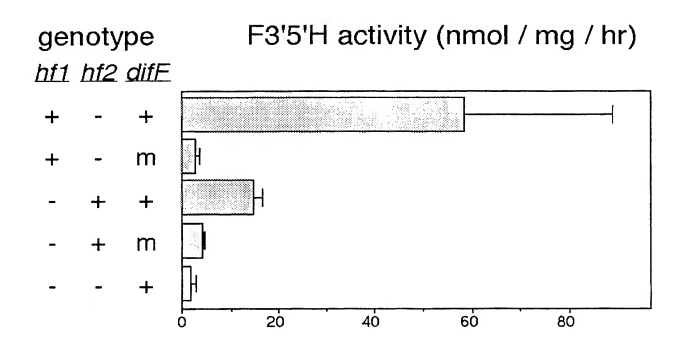


Figure 4d

	AAA	TGG.	ACA	AAC?	AAA	GAG	TGI	TLL	ACAC	TTT	rcrc	PAAC	3TC(3CAC	3AA(CAC	AAG	TCA	AAG	CAA	GAT	TGC	TGG	ATT	AAATGGACAAACAAAGAGTGTTTACACTTTTCTCAAGTCGCAGAACACAAGTCAAAGCAAGATTGCTGGATTATCA		75
	Σ	Q	×	Ø	었	>	[<u>T</u> 4		T I	1	S	C	Q V A		四	H	×	Ø	K O D C W	œ	Ω	ບ		н	п	(2)	(25)
	TCA	ATGGC x2324	GCA 24	GA G	IAG	TAG	ATG	TA.	ACAZ	\AG	[TC]	rtge	3AA(3AA(CAT(CCT	3GA	GGA	GAA	GAA	GTG	TTG	ATT	GAA	TCAATGGCAGAGTAGTAGATGTAACAAGTTCTTGGAAGAACATCCTGGAGGAGAAGAAGTGTTGATTGA		150
	Z	ט	24	>	>	Д	>		H	×	E4	回	(r)	团	н	O.	r U	_O	ы Б Б Б		>	V L I		臼	S A	3)	(20)
S	CAG	GAA	AGG	ATG(CAA	CTA	AAG	:AG	[TTC	PAAC	3AT?	\TT(3GA(ZAT	AGT.	AAA	GCT	ညည	'AAG	AAC	TTG	CII	TTC	'AAZ	CAGGAAAGGATGCAACTAAAGAGTTTCAAGATATTGGACATAGTAAAGCTGCCAAGAACTTGCTTTTCAAATACC	.,	225
JBS'	ŋ	×	А	G K D A	H	¥,	国		C ₄	7 I) ⊔	רי	##	S	₩	Æ	Ø	Q D I G H S K A A K N L L F K Y	Z	ы	ы	ᄄ	×	o X	S	(75)
TITUTE	AAA	TTG	GAT	ATC.	IIC	AAG	GTJ	X2	ACAAAC X2325) [C	I'CA(3AT(3AT	ICT	GAA)	CTT	GAA	CTC	AAC	TI	GTC	PAC	GAI	TCC	AAATTGGATATCTTCAAGGT <u>TACAAAG</u> CCTCAGATGATTCTGAACTTGAACTCAACTTAGTCACTGATTCCATCA X2325	.,	300
SHI	H	r U	> 4	н	Ø	Ö	¥		K 7	€	S	-	D D		四	н	凶	ı	LNLVTDSI	П	>	H	Q	ß	I	(1((100)
EET (AAC	CAA	ATA	AGG	CCA	AAG	JAA1	ATG2	4AA(3CT.	rat(3TT.	ATC	AAA	GAA	GAT	CCI	'AAG	CC	AAG	TA	CTC	3AC	AAGAACCAAATAAGGCCAAAGAAATGAAAGCTTATGTTATCAAAGAAGATCCTAAGCCAAAGTATCTGACTTTTG	'n	375)
(\mathbf{RU})	闰	Д	Z	×	Ą	X	田		X	 Y	a!	ы	>	H	×	[L]	Д	Д	KAYVIKEDPKPKYLT	വ	×	₩	ļ	EH	[τ ₄	(1;	(125)
LE 26		AGT	ACT	TAT	IGC	CCI	TCI	TGC	3CTC	3CT(300	LTC	TAC	CIC	TAT	TAT	ညည	TAT	CTC	ACT	GGP	D D	CTC	Š	TTGAGTACTTATTGCCCTTCTTGGCTGCCTTCTACCTCTATTATCGCTATCTCACTGGAGCTCTCCAGTTTT	•	45(
5)		>	<u></u>	EYLL	Д	ľų -	 C-	T 7	A A		Æ	[zr	 >₁		≯₁	> 1	ρú	×	AAFYLYYRYLTG	E	_O	A	A L Q	ø	* [I4	(1)	(149)
	GAG	GIC	AGA	GAA(CAA	AGG	ATT	PAC	4CT2	ACA:	rga.	lta.	LTG	TCA	GTA'	TAT	TCT	CAC	TGG	AGC	TAT	55	'AT	GTJ	GAGCTCAGAGAACAAAGGATTACACTACATGATTATTGTCAGTATATTCTCACTGGAGCTATCGCATTGTTTGAA		525
	CCT	TAG	AAG	ATA(CTT	GGI	GAI	TTC	rgg2	AAA	AGT(3TT	TTC	TTT.	ATT	TAT	TTT	'AA1	CTI	CA	AGA	AAC	CTC	3GA(CCTTAGAAGATACTTGGTGATTCTGGAAAAGTGTTTTTTTT		900
	TTG	ATT	GTT	ATT(CIT	GCI	TG	TC	ATT	ICA	3AA(CIA	CTG	AAC.	AGT	TTT	CCA	ACC	CAC	TT	GAG	CAC	AAC	ŢŢ	TIGATIGITATICITIGCTIGITCATITICAGAACTACIGAACAGITITICCAACCCACTITGAGCACAACTCTITAT		675
																											1

Figure 5

TCTAAAAAAAAAAAAAAAAAAAAAAAAAAA

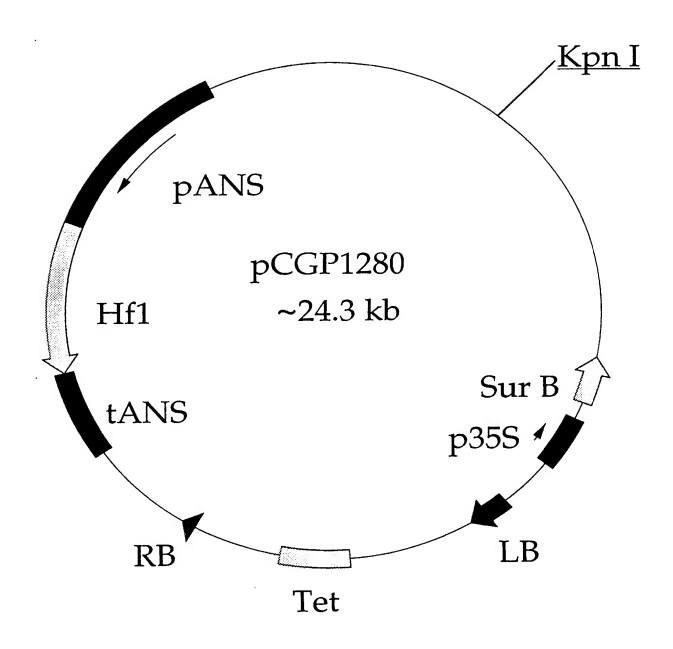


Figure 6a **SUBSTITUTE SHEET (RULE 26)**

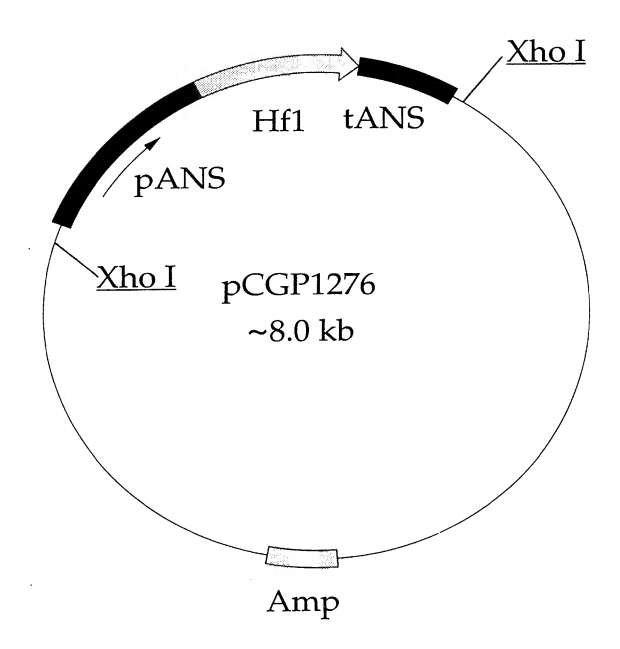


Figure 6b **SUBSTITUTE SHEET (RULE 26)**

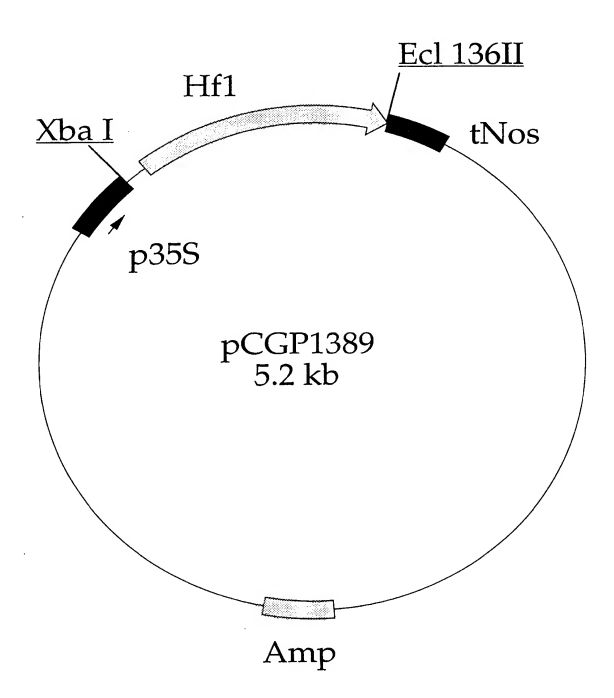


Figure 6c **SUBSTITUTE SHEET (RULE 26)**

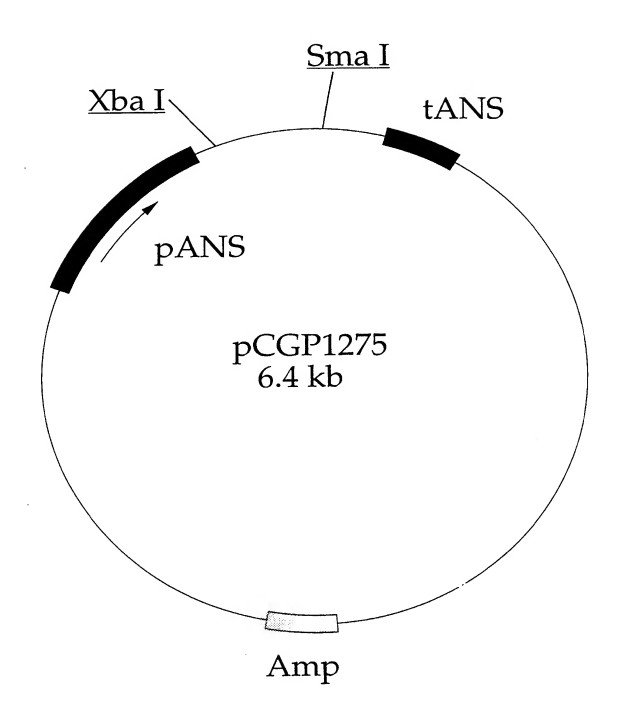


Figure 6d **SUBSTITUTE SHEET (RULE 26)**

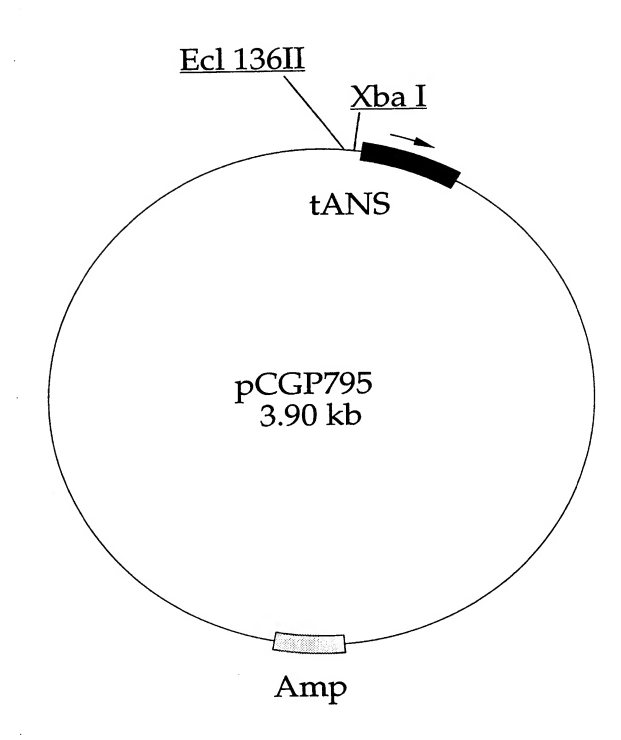


Figure 6e **SUBSTITUTE SHEET (RULE 26)**

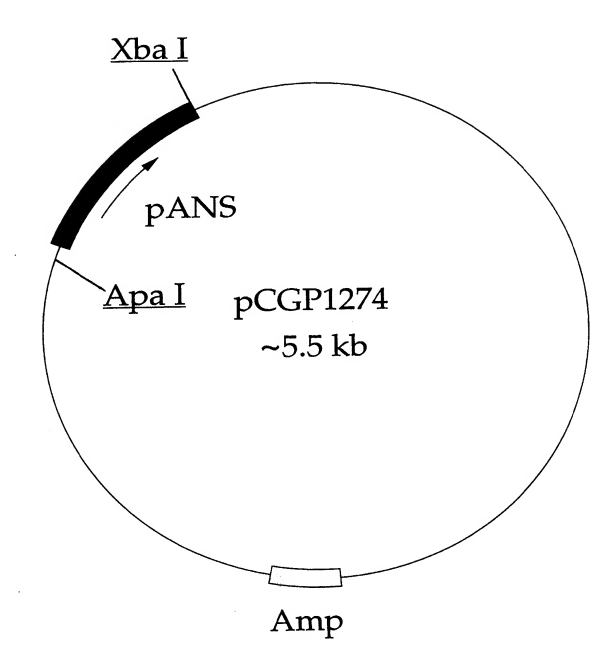


Figure 6f **SUBSTITUTE SHEET (RULE 26)**

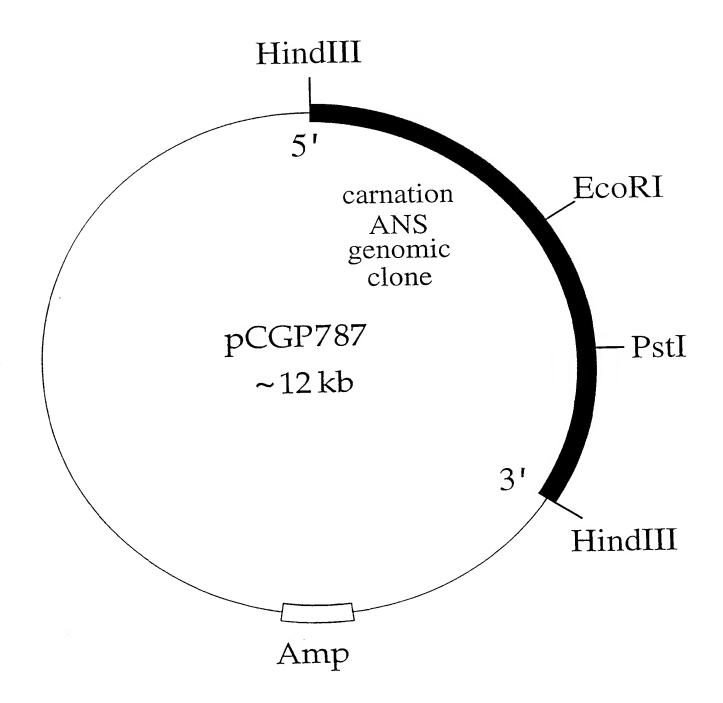
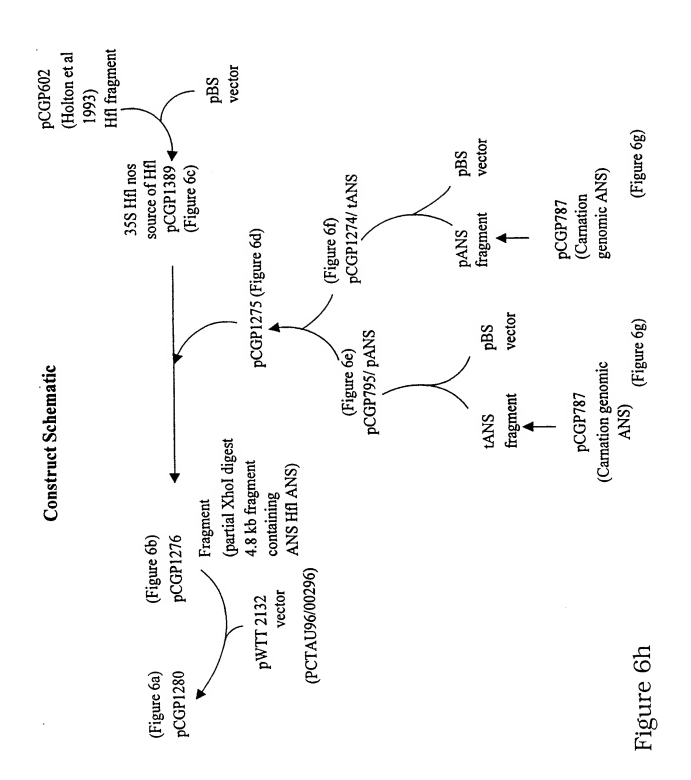


Figure 6g SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)

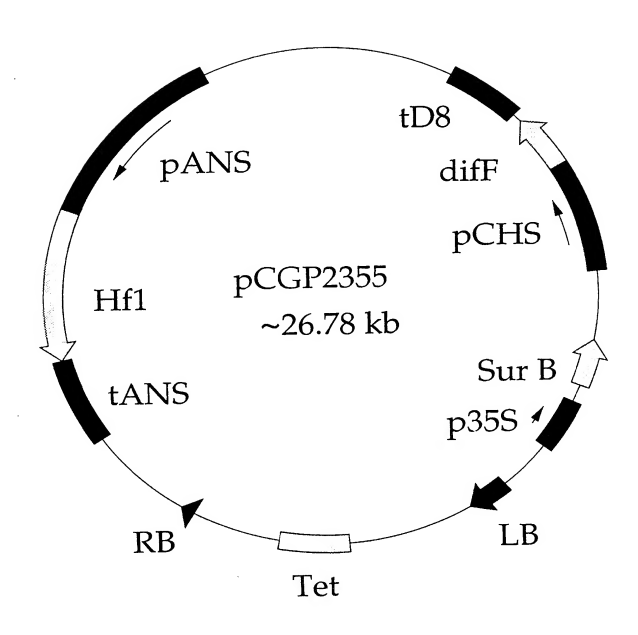


Figure 7a **SUBSTITUTE SHEET (RULE 26)**

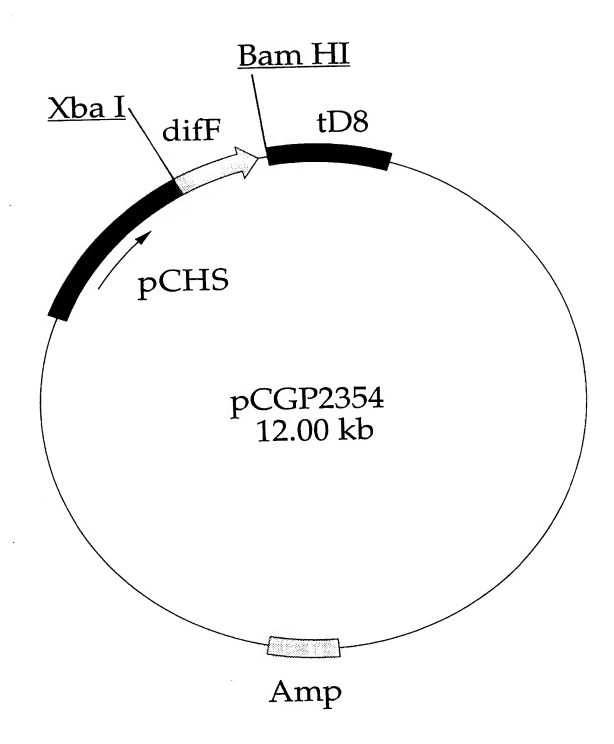


Figure 7b SUBSTITUTE SHEET (RULE 26)

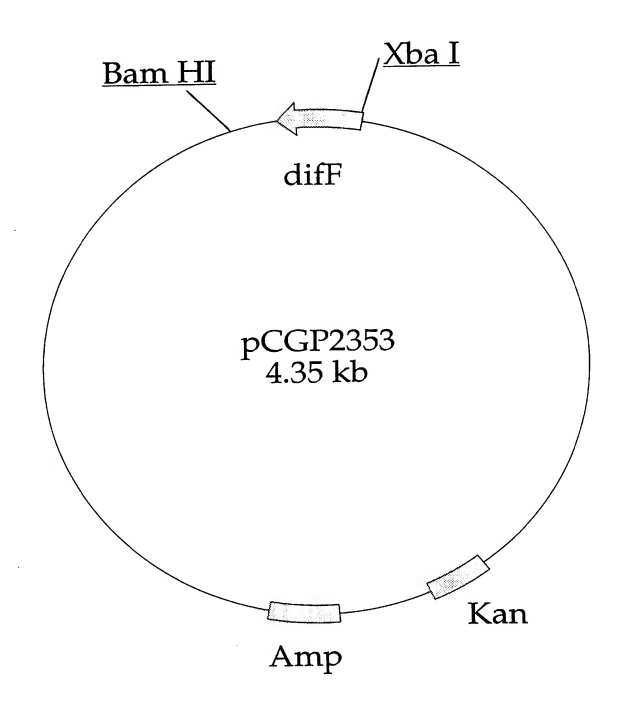


Figure 7c SUBSTITUTE SHEET (RULE 26)

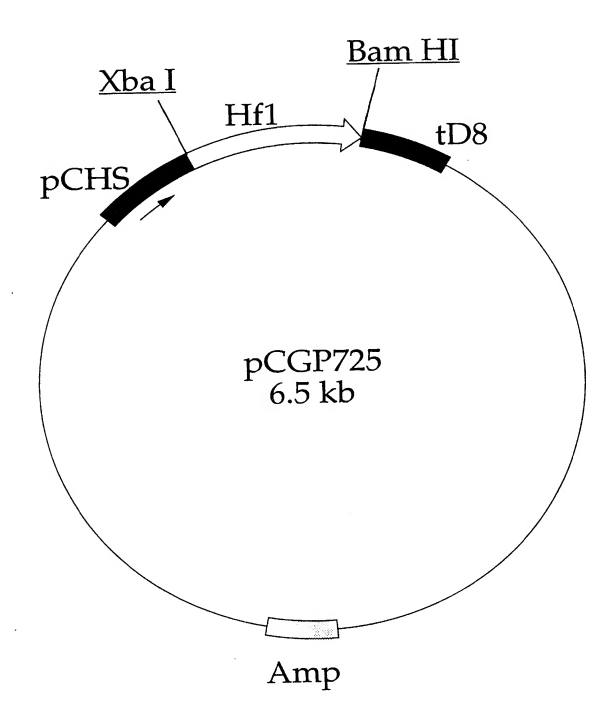


Figure 7d **SUBSTITUTE SHEET (RULE 26)**

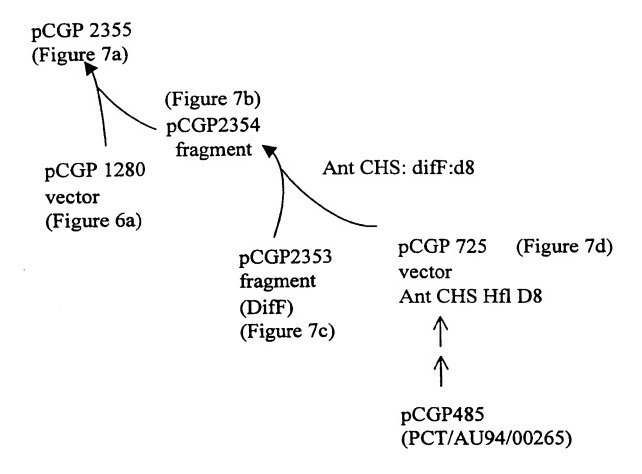


Figure 7e

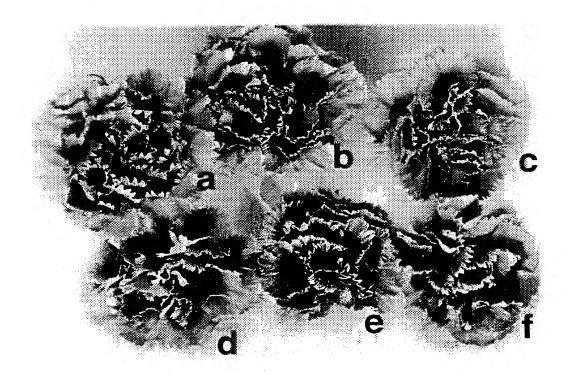


Figure 8

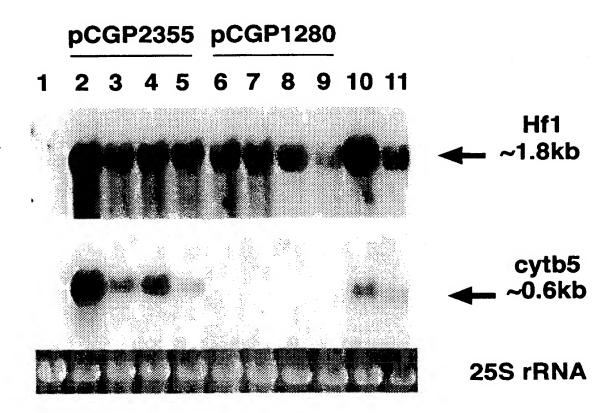


Figure 9 SUBSTITUTE SHEET (RULE 26)

- 1 -

SEQUENCE LISTING

<110> INTERNATIONAL FLOWER DEVELOPMENT AND VRIJE UNIVERSITEIT <120> A PLANT REGULATORY GENE <130> 2204892/EJH <140> <141> <150> PP5268 <151> 1998-08-14 <160> 5 <170> PatentIn Ver. 2.0 <210> 1 <211> 707 <212> DNA <213> Petunia sp. <220> <221> CDS <222> (3)..(449) <400> 1 aa atg gac aaa caa aga gtg ttt aca ctt tct caa gtc gca gaa cac 47 Met Asp Lys Gln Arg Val Phe Thr Leu Ser Gln Val Ala Glu His 15 10

95	gc aga gta gta gat	aat	atc	atc	att	tgg	tgc	gat	caa	aag	tca	aag
	ly Arg Val Val Asp	Asn	Ile	Ile	Ile	Trp	Cys	Asp	Gln	Lys	Ser	Гуs
	30		25					20				
143	aa gaa gtg ttg att	gga g	gga	cct	cat	gaa	gaa	ttg	ttc	aag	aca	gta
	lu Glu Val Leu Ile	Gly	Gly	Pro	His	Glu	Glu	Leu	Phe	Lys	Thr	Val
	45			40					35			
191	aa gat att gga cat	ttt c	gag	aaa	act	gca	gat	aag	gga	gca	tca	gaa
	In Asp Ile Gly His	n Phe	Glu	Lys	Thr	Ala	Asp	Lys	Gly	Ala	Ser	Glu
	60				55					50		
239	aa att gga tat ctt	tac c	aaa	ttc	ctt	ttg	aac	aag	gcc	gct	aaa	agt
	Sln Ile Gly Tyr Leu	s Tyr	Lys	Phe	Leu	Leu	Asn	Lys	Ala	Ala	Lys	Ser
	75					70					65	
287	aa ctc aac tta gtc	ctt (gaa	tct	gat	gat	tca	gcc	aaa	tac	ggt	caa
	Blu Leu Asn Leu Val	ı Leu	Glu	Ser	Asp	Asp	Ser	Ala	Lys	Tyr	Gly	Gln
	95	90					85					80
335	aa atg aaa gct tat	aaa	gcc	aag	aat	cca	gaa	aaa	atc	tcc	gat	act
	Glu Met Lys Ala Tyr	a Lys	a Ala	Lys	Asn	Pro	Glu	Lys	Ile	Ser	Asp	Thr
	110	5	105				ı	100				
						-						
383	ct ttt gtt gag tac	ctg	tat	aag	cca	aag	cct	gat	gaa	aaa	ato	gtt
	Thr Phe Val Glu Tyr	r Leu	з Туз	Lys	Pro	Lys	Pro	Asp	Glu	Lys	Ile	Val
	125		0	120				•	115			
431	at tat cgc tat ctc	ctc	tac	ttc	gcc	gct	gct	ttg	ttc	ccc	tto	tta

PCT/GB99/02676 WO 00/09720

- 3 -

Leu Leu Pro Phe Leu Ala Ala Phe Tyr Leu Tyr Tyr Arg Tyr Leu 140 135 130

479 act gga gct ctc cag ttt tgagctcaga gaacaaagga ttacactaca Thr Gly Ala Leu Gln Phe

tgattattgt cagtatattc tcactggagc tatcgcattg tttgaacctt agaagatact 539 tggtgattct ggaaaagtgt tttctttatt tattttaatc ttcaaagaaa gctggagtta 599 cttgattgtt attcttgctt gttcatttca gaactactga acagttttcc aacccacttt 659 707

<210> 2

<211> 149

145

<212> PRT

<213> Petunia sp.

<400> 2

Met Asp Lys Gln Arg Val Phe Thr Leu Ser Gln Val Ala Glu His Lys 15 1 5 10

Ser Lys Gln Asp Cys Trp Ile Ile Ile Asn Gly Arg Val Val Asp Val 25 30 20

Thr Lys Phe Leu Glu Glu His Pro Gly Gly Glu Glu Val Leu Ile Glu 40 45 35

-4-

Ser Ala Gly Lys Asp Ala Thr Lys Glu Phe Gln Asp Ile Gly His Ser 60 55 50 Lys Ala Ala Lys Asn Leu Leu Phe Lys Tyr Gln Ile Gly Tyr Leu Gln 75 80 70 65 Gly Tyr Lys Ala Ser Asp Asp Ser Glu Leu Glu Leu Asn Leu Val Thr 90 95 85 Asp Ser Ile Lys Glu Pro Asn Lys Ala Lys Glu Met Lys Ala Tyr Val 110 105 100 Ile Lys Glu Asp Pro Lys Pro Lys Tyr Leu Thr Phe Val Glu Tyr Leu 125 115 120 Leu Pro Phe Leu Ala Ala Phe Tyr Leu Tyr Tyr Arg Tyr Leu Thr 140 135 130 Gly Ala Leu Gln Phe 145 <210> 3 <211> 149 <212> PRT <213> Petunia sp. <400> 3 Met Asp Lys Gln Arg Val Phe Thr Leu Ser Gln Val Ala Glu His Lys 10 15

Ser Lys Gln Asp Cys Trp Ile Ile Ile Asn Gly Arg Val Val Asp Val
20 25 30

Thr Lys Phe Leu Glu Glu His Pro Gly Gly Glu Glu Val Leu Ile Glu
35 40 45

Ser Ala Gly Lys Asp Ala Thr Lys Glu Phe Gln Asp Ile Gly His Ser
50 55 60

Lys Ala Ala Lys Asn Leu Leu Phe Lys Tyr Gln Ile Gly Tyr Leu Gln 65 70 75 80

Gly Tyr Lys Ala Ser Asp Asp Ser Glu Leu Glu Leu Asn Leu Val Thr

85 90 95

Asp Ser Ile Lys Glu Pro Asn Lys Ala Lys Glu Met Lys Ala Tyr Val

Ile Lys Glu Asp Pro Lys Pro Lys Tyr Leu Thr Phe Val Glu Tyr Leu
115 120 125

Leu Pro Phe Leu Ala Ala Phe Tyr Leu Tyr Tyr Arg Tyr Leu Thr

Gly Ala Leu Gln Phe

145

<210> 4

<211> 2552

PCT/GB99/02676

- 6 -

<212> DNA

<213> carnation

<400> 4

gaattcactc gaaatgatac tgtgagtctg cgataggctc gttttgaggc ggaaattatg 60 attttacgac gtgatcaatt gagcatgact taaatttgcg tcttctcagt cgtcgttgca 120 ttgcaatttt tagtattttc aggtgctctg aaagtgttta gtacatcgtt tttaaaatgg 180 atatettttt gttetggteg acttaetett egetttttaa tgeagaegtg eeegttattg 240 ctacgtgtat tcacaaaggt atgaccgtgt tctgtagcgt ctaatgataa tatatgaagt 300 cgaggttgca tttgtactag tccgataata attagtatcg ttttcatact gatactagat 360 cggaggtcac catacccgtg aagatttttc tgtgagagga aaagaaccca aggacgaggt 420 tcaaatctac acatggaaag acgccacgct tcgtgagtta actgaccttg tatgtcgcca 480 togottagog tagogotgaa catogittito accotgotoo atocatoaac catitatigg 540 tottatacat gtgtgattgc gttgttotta catttaggtg aaagacgttt ctccagctgc 600 taggagtcga gatgcgaaat tgtcgtttgc gactgtatac cttgatagaa atggatgcat 660 gcaagtaaag aaggtatett etaatteate tttegtagag acatagegtg aatttggaeg 720 gggtctttgg tttgagaaag ataacagctt tacgtatttt tgtagatggg tgaaaccttt 780

tcaaatccgt ataagcgtaa agacgacaac tgggctttag gggacacatt ctttcaggta 840 taattgatgc gactaacaat agtctccact gatcatattc tactcttcta cgttcgatac 900 tgactgtttc tggttatttg gtagacagga gattatttgg acgtagcaat tcagtagcgt 960 agagatgttt ccacacgtgt tatcgtaaaa gaagcaagat aagcctaatg cctagggtgg 1020 tggtatgact teegttgett ategategtg ettgtaagta attteegtet tatettttee 1080 tgttatataa agttaatett etetaggaet tteatgaace ttgtttgtgt atttatttet 1140 cgatcaacat gatagagcta gtttttaagc aacgtatact agtagtctat tggaagttaa 1200 gacacggttc ttaaaaaggt acgatccaag tgaagcatgt tagatatgac actttcttct 1260 agggacgact ctcgtatgcc acccgacttt ttcaattttt tttgtgaatg ttagatgtgt 1320 gtatataatg catccgaaag atgtctcaac gaacaaatga gccacctact tcgatcactc 1380 gctatcaatg ttattaatgc cttgttgatt ttaatagttg atcaataata gtaaaatcta 1440 ttcaagggta tagtctcccg ttcacactca tcggggttac actagcgagc tccattaatc 1500 ggtgccttaa tcgagacgct aagaactata ccatgaccta gtcagcgcca tgggactgat 1560 gtaggccaca caatctcgat gatccgaaaa cgctagagtt caagacctag ttcgagacca 1620 tggtcacggt ttcaaccgcg atatctcaac aatgcaattt ttttcgagac tagacagacg 1680 aataagtett gtgtacgatg ggtagetagt gaattaaagg taateaettt aetegtgtte 1740

acaagaagac cattcatatg aacatttcgt gttctcagac gagctccttt cgctagtttg 1800 gtaaacttgt ttggtgttac ctggcttatc atagcccact caatttggcg aaaacataac 1860 aattgtttca catgcgaagg tctacacacc catactcgtg ataaaaacgg ttgcatttat 1920 tattattgcc tttcgagcaa tttacgttgt ttgaagtttg tgtaaaaaca aaatcgatct 1980 ataattactg actggaacgg taatatcaaa actgttagga cctgttcttt tcggcttata 2040 tgaactgaac ttataagggc tgaaattaag ctgtaaagaa caggttetta tttegeetga 2160 tttgaatcac taaagttgga tgaagataga ggcgagaaac gctccccttt ccacagtttg 2220 actctactcg aatccgacca aaattggatg taaattagaa gtgagcattc ctcatgccaa 2280 ccaccgttct tggtcgtgaa aacatgattt gtagctgctg tatacttccc aaacccgtga 2340 taatctgcca cacttccaac acttaattga tttttatcaa aattcctaca gtttttttggt 2400 tttactccta aactatgctg tctttttaca agttgttaca cctttgtcaa caactttatg 2460 ctttattata tttcttatat aaagaccata taacttccct acactaatgc cacaacactt 2520 2552 aaaagcatac aacacaaacc tcataatcta ga

<210> 5

<211> 2552

<212> DNA

<213> carnation

<400> 5

tctagattat gaggtttgtg ttgtatgctt ttaagtgttg tggcattagt gtagggaagt 60 tatatggtct ttatataaga aatataataa agcataaagt tgttgacaaa ggtgtaacaa 120 cttgtaaaaa gacagcatag tttaggagta aaaccaaaaa actgtaggaa ttttgataaa 180 aatcaattaa gtgttggaag tgtggcagat tatcacgggt ttgggaagta tacagcagct 240 acaaatcatg ttttcacgac caagaacggt ggttggcatg aggaatgctc acttctaatt 300 tacatccaat tttggtcgga ttcgagtaga gtcaaactgt ggaaagggga gcgtttctcg 360 cctctatctt catccaactt tagtgattca aatcaggcga aataagaacc tgttctttac 420 agettaattt cageeettat aagtteagtt caactettat aagtteagtt cageteaget 480 cctataagta cagcttttat aagttcagct cttataagcc gaaaagaaca ggtcctaaca 540 gttttgatat taccgttcca gtcagtaatt atagatcgat tttgttttta cacaaacttc 600 aaacaacgta aattgctcga aaggcaataa taataaatgc aaccgttttt atcacgagta 660 tgggtgtgta gaccttcgca tgtgaaacaa ttgttatgtt ttcgccaaat tgagtgggct 720 atgataagcc aggtaacacc aaacaagttt accaaactag cgaaaggagc tcgtctgaga 780

acacgaaatg ttcatatgaa tggtcttctt gtgaacacga gtaaagtgat tacctttaat 840 tcactagcta cccatcgtac acaagactta ttcgtctgtc tagtctcgaa aaaaattgca 900 ttgttgagat atcgcggttg aaaccgtgac catggtctcg aactaggtct tgaactctag 960 cgttttcgga tcatcgagat tgtgtggcct acatcagtcc catggcgctg actaggtcat 1020 ggtatagttc ttagcgtctc gattaaggca ccgattaatg gagctcgcta gtgtaacccc 1080 gatgagtgtg aacgggagac tataccettg aatagatttt actattattg atcaactatt 1140 aaaatcaaca aggcattaat aacattgata gcgagtgatc gaagtaggtg gctcatttgt 1200 tcgttgagac atctttcgga tgcattatat acacacatct aacattcaca aaaaaaattg 1260 aaaaagtcgg gtggcatacg agagtcgtcc ctagaagaaa gtgtcatatc taacatgctt 1320 cacttggatc gtaccttttt aagaaccgtg tcttaacttc caatagacta ctagtatacg 1380 ttgcttaaaa actagctcta tcatgttgat cgagaaataa atacacaaac aaggttcatg 1440 aaagtcctag agaagattaa ctttatataa caggaaaaga taagacggaa attacttaca 1500 agcacgatcg ataagcaacg gaagtcatac caccacccta ggcattaggc ttatcttgct 1560 tettttacga taacaegtgt ggaaacatet etaegetaet gaattgetae gtecaaataa 1620 tctcctgtct accaaataac cagaaacagt cagtatcgaa cgtagaagag tagaatatga 1680

tcagtggaga ctattgttag tcgcatcaat tatacctgaa agaatgtgtc ccctaaagcc 1740 cagttgtcgt ctttacgctt atacggattt gaaaaggttt cacccatcta caaaaatacg 1800 taaagetgtt atetttetea aaccaaagae eeegteeaaa tteaegetat gtetetaega 1860 aagatgaatt agaagatacc ttctttactt gcatgcatcc atttctatca aggtatacag 1920 togoaaacga caatttogoa totogactoo tagoagotgg agaaacgtot ttoacotaaa 1980 tgtaagaaca acgcaatcac acatgtataa gaccaataaa tggttgatgg atggagcagg 2040 gtgaaaacga tgttcagcgc tacgctaagc gatggcgaca tacaaggtca gttaactcac 2100 gaagcgtggc gtctttccat gtgtagattt gaacctcgtc cttgggttct tttcctctca 2160 cagaaaaatc ttcacgggta tggtgacctc cgatctagta tcagtatgaa aacgatacta 2220 attattatcg gactagtaca aatgcaacct cgacttcata tattatcatt agacgctaca 2280 gaacacggtc atacctttgt gaatacacgt agcaataacg ggcacgtctg cattaaaaag 2340 cgaagagtaa gtcgaccaga acaaaaagat atccatttta aaaacgatgt actaaacact 2400 ttcagagcac ctgaaaatac taaaaattgc aatgcaacga cgactgagaa gacgcaaatt 2460 taagtcatgc tcaattgatc acgtcgtaaa atcataattt ccgcctcaaa acgagcctat 2520 cgcagactca cagtatcatt tcgagtgaat tc